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Kruppel-like Factor 15 Is a Critical Regulator of Cardiac Lipid Metabolism*^S

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Background: Metabolic homeostasis is central to normal cardiac function. The molecular mechanisms underlying metabolic plasticity in the heart remain poorly understood.

Results: Kruppel-like factor 15 (KLF15) is a direct and independent regulator of myocardial lipid flux.

Conclusion: KLF15 is a core component of the transcriptional circuitry that governs cardiac metabolism.

Significance: This work is the first to implicate the KLF transcription factor family in cardiac metabolism.

The mammalian heart, the body's largest energy consumer, has evolved robust mechanisms to tightly couple fuel supply with energy demand across a wide range of physiologic and pathophysiologic states, yet, when compared with other organs, relatively little is known about the molecular machinery that directly governs metabolic plasticity in the heart. Although previous studies have defined Kruppel-like factor 15 (KLF15) as a transcriptional repressor of pathologic cardiac hypertrophy, a direct role for the KLF family in cardiac metabolism has not been previously established. We show in human heart samples that KLF15 is induced after birth and reduced in heart failure, a myocardial expression pattern that parallels reliance on lipid oxidation. Isolated working heart studies and unbiased transcriptomic profiling in Klf15-deficient hearts demonstrate that KLF15 is an essential regulator of lipid flux and metabolic homeostasis in the adult myocardium. An important mechanism by which KLF15 regulates its direct transcriptional targets is via interaction with p300 and recruitment of this critical co-activa-

tor to promoters. This study establishes KLF15 as a key regulator of myocardial lipid utilization and is the first to implicate the KLF transcription factor family in cardiac metabolism.

The working mammalian heart requires enormous amounts of energy to maintain contractile function and blood flow. Under physiologic conditions, the majority this energy production is derived from oxidative catabolism of fatty acids (FAs)³ (~70%), with the remainder being derived from glucose (~20%) and other substrates, such as lactate and ketones (~10%) (1). In order to maintain contractile performance under a wide range of physiologic and pathophysiologic settings, the myocardium has evolved critical molecular mechanisms that match fuel supply with energy demand and nutrient availability (2–4).

A major form of the heart's metabolic plasticity is its ability to augment lipid utilization as a means to guard against energy depletion and circulatory failure during periods of transient carbohydrate deprivation (*e.g.* fasting) or heightened energy needs (*e.g.* exercise, postnatal growth). Defects in specific genes critical for efficient myocardial lipid utilization are known to cause metabolic cardiomyopathies in both animal models and humans (5). In addition, loss of metabolic plasticity has been observed in common forms of human heart failure (5–8), a leading cause of hospitalization and death worldwide (5). Importantly, recent studies strongly support the contention

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^S This article contains supplemental Table 1.

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³ The abbreviations used are: FA, fatty acid; TG, triglyceride; PPAR, peroxisome proliferator-activated receptor; ERR, estrogen-related receptor; KLF, Kruppel-like factor; NRVM, Neonatal rat ventricular myocyte(s); LVAD, left ventricular assisted device; qPCR, quantitative PCR; IP, immunoprecipitation; IWH, isolated working heart.

that dysregulated lipid utilization and energy homeostasis play a causal role in heart failure progression (9, 10). As recently highlighted by Goldberg et al. (2), although the heart is far and away the most energy-requiring organ in the body, studies of cardiac lipid metabolism, especially *in vivo* are relatively scarce compared with investigations in adipose tissue, liver, or skeletal muscle. A better understanding of the fundamental mechanisms governing metabolic plasticity in the myocardium may therefore provide key insights into heart failure pathogenesis and pave the way for novel therapeutic approaches for this common and lethal condition.

Efficient myocardial lipid flux involves coordination of several key enzymatic steps, including sarcolemmal FA uptake and activation, maintenance of the intracellular triglyceride (TG) pool, acyl-CoA import into the mitochondrial matrix, β-oxidation, TCA cycle, and oxidative phosphorylation (1, 2, 5, 11). The enzymatic machinery regulating each of these steps is under robust allosteric and transcriptional control in a manner that tightly couples lipid flux with energy demand and nutrient availability (12, 13). Despite the fundamental importance of metabolic plasticity in the heart, there is surprisingly little known about the transcriptional pathways that directly regulate myocardial substrate flux. When compared with the vast body of work that has defined gene-regulatory networks governing myocyte hypertrophy (14), the identification of transcription factors that are bona fide and direct regulators of myocardial nutrient flux is limited to a few members of the nuclear receptor superfamily (PPARs and ERRs) and the forkhead box factor FoxO1 (12, 15-17).

Kruppel-like factors (KLFs) are DNA-binding transcriptional regulators that contain three conserved zinc fingers within the carboxyl terminus that bind a putative consensus 5'-C(A/T)CCC-3' motif in the promoters and enhancers of various genes (18, 19). The amino terminus is involved in transcriptional activation and repression as well as protein-protein interaction. Previous studies from our group have defined KLF15 as a transcriptional repressor of pathologic cardiac hypertrophy (20–22). In contrast, the broad transcriptional programs directly regulated by KLF15 in the heart remain unknown.

Initial insights linking the KLF transcription factor family to metabolic homeostasis were gleaned from our studies defining KLF15 as a regulator of adipogenesis (23). Subsequent studies in liver and skeletal muscle in the context of circadian cycles, fasting, and exercise now identify KLF15 as a central component of the transcriptional circuitry that regulates physiologic flux of all three major nutrient classes (glucose, lipids, and amino acids) (24–26). Despite these important observations, no role for any member of the KLF family in cardiac metabolism has been elucidated.

Here, we demonstrate that Kruppel-like factor 15 (KLF15) is a direct and independent regulator of myocardial lipid flux *in vivo* through a molecular mechanism involving recruitment of the chromatin remodeling enzyme p300 to target promoters. This study introduces an entirely new class of transcriptional regulators (KLFs) to the core molecular circuitry that governs myocardial metabolism.

EXPERIMENTAL PROCEDURES

Animal Models—All protocols concerning animal use were approved by the Institutional Animal Care and Use Committee at Case Western Reserve University and conducted in strict accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. *Klf15^{-/-}* mice have been described previously (23). Studies were performed with age-and sex-matched littermate controls (12–14 weeks old, male, pure C57Bl/6 background). Mice were housed in a temperature- and humidity-controlled barrier facility with a 12-h light/ dark cycle and *ad libitum* access to water and standard laboratory rodent chow (Laboratory diet P3000; 4.5% fat by weight, 14% kcal).

Cell Culture—Neonatal rat ventricular myocytes (NRVM) were isolated from 2-day-old rat pups and isolated and maintained under standard conditions as described previously (20, 21, 27). Isolated NRVM were cultured for 24-48 h under quiescent conditions by the inclusion of serum-free DMEM supplemented with insulin, transferrin, and selenium prior to experiments. C2C12 myoblasts were purchased from ATCC (Manassas, VA) and grown in DMEM supplemented with 10% FBS. C2C12 myoblasts were passaged prior to reaching 60% confluence to maintain cells in the undifferentiated state. All studies used low passage C2C12 (passages 3-6). $P300^{-/-}$ mouse embryonic fibroblasts (28) were cultured in DMEM supplemented with 10% FBS as described previously (29).

Plasmids and Adenoviruses—Expression plasmids containing the mouse KLF15 cDNA have been described previously (20). The –1.2 kb promoter region of mouse *Slc27a1* (*Fatp1*) was PCR-amplified from purified genomic DNA (Promega) and subcloned into pGL3-basic (Promega). The *Pdk4* promoter-luciferase construct was a gift from D. Kelly (30). For shRNA against mouse/rat *Klf15*, the hairpin sequence 5'-GCGGTAA-GATGTACATCAA<u>ACGTGTGCTGTCCGT</u>TTGGTGTAC-ATCTTGCTGC-3' (loop sequence is underlined) was subcloned into the pEQ adenoviral shRNA vector (Welgen, Inc.). Recombinant adenoviruses for sh-*Klf15* and sh-scrambled were amplified and purified by Welgen, Inc.

Isolated Working Heart—The isolated mouse working heart perfusion was performed at the Washington University Metabolic Core as described previously (31, 32). In brief, hearts were excised and placed in ice-cold Krebs-Henseleit solution. Retrograde perfusion of the aorta was first performed via the Langendorff method followed by left atrial cannulation. Working mode was maintained in oxygenated buffer containing 0.4 mM palmitate, 5.0 mM glucose, and 100 microunits insulin/ml with a preload pressure of 11.5 mm Hg and afterload pressure of 50 mm Hg. Palmitate and glucose oxidation rates were determined using radiolabeled [³H]palmitate and [U-¹⁴C]glucose, respectively. Samples were collected every 10 min, and ³H₂O and ¹⁴CO₂ radioactivity was counted to determine palmitate and glucose oxidation rates, respectively. In addition, functional measurements such as cardiac output, peak systolic pressure, and heart rate were measured as described previously (31).

Human Left Ventricular Samples—Human left ventricular RNA samples were obtained as described previously (6). In brief, myocardial specimens were collected before and after left





ventricular assisted device (LVAD) implantation and explantation as a bridge to transplantation for end-stage HF patients. Control heart samples were obtained from non-failing hearts as described previously (6). Fetal left ventricular samples were obtained at the time of elective termination immediately following tissue extraction. The use of all human samples was approved by the Institutional Review Board of Columbia University and Case Western Reserve University (IRB-AAAE7393).

Measurement of Plasma Parameters and Tissue Lipids— Hearts and plasma were rapidly harvested and flash frozen from WT and KO mice. Lipids were extracted from ventricles by the Folch method. Cardiac FA and TG content was quantified by thin layer chromatography by the National Institutes of Health-Mouse Metabolic Phenotyping Core/Lipid Analytic Core at Vanderbilt University. Plasma free fatty acids and TGs were quantified by the Vanderbilt National Institutes of Health Mouse Metabolic Phenotyping Core.

Mitochondrial Physiology and Electron Microscopy—Myocardia of four individual WT or KO mice were pooled for each experiment. The data presented are from four independent experiments. Fresh myocardial subsarcolemmal mitochondria were isolated as described previously (33). Mitochondrial protein concentration, citrate synthase activity, and oxygen consumption (using a Clark-type oxygen electrode at 30 °C) were assayed as described previously (33, 34). Mouse heart tissue was freshly dissected and rapidly processed for transmission electron microscopy as described previously (35).

Mitochondrial Genome Quantification—Total tissue DNA was prepared from hearts of WT versus KO mice using the QIAamp DNA minikit (Qiagen) according to the manufacturer's instructions. SYBR Green qPCR of purified DNA was performed using primers specific for the mouse mitochondrial Mt-Cox2 gene. Values were normalized to the nucleus-encoded mouse Rplp0 gene using the $\Delta\Delta Ct$ method. Primer sequences are available upon request.

Chromatin Immunoprecipitation-Chromatin isolated from 15 million NRVM were used for each immunoprecipitation. NRVM were infected with empty virus or Ad-KLF15-FLAG, fixed with 1.1% formaldehyde for 10 min, and quenched with 0.125 M glycine for 5 min. Chromatin was extracted and sonicated using a BioRuptor (Diagnode, Sparta, NJ). The sonicated chromatin was immunoprecipitated with 5 μ g of anti-FLAG(M2) antibodies (Sigma) bound to Dynabeads (Invitrogen) followed by extensive washing and elution. For heart tissue, chromatin isolated from two hearts was pooled for a single immunoprecipitation. Tissue was finely minced in PBS supplemented with protease inhibitors, fixed with 1.1% formaldehyde (10 min), and quenched with 0.125 M glycine (5 min). Tissues were disrupted using a Dounce homogenizer followed by chromatin extraction and sonication with a Bioruptor. Approximately 500 µg of chromatin was used in each immunoprecipitation with 5 μ g of anti-p300 antibody (C-20, Santa Cruz Biotechnology, Inc.) bound to Dynabeads followed by extensive washing and elution. Chromatin was then reverse cross-linked followed by purification of genomic DNA. Target and non-target regions of genomic DNA were amplified by qPCR in both the immunoprecipitates and input samples. For NRVM samples, the rat Alb1 locus was used as a non-target control. For

mouse heart tissue samples, the 28 S genomic locus was used as a non-target control. Relative abundance in immunoprecipitates was expressed as the percentage of abundance in input samples as described previously (26, 36). ChIP primer sequences are available upon request.

RNA Extraction and qPCR—Tissue samples were disrupted/ homogenized in PureZOLTM (Bio-Rad) in a Tissue-lyzer (Qiagen), using stainless steel beads (30 Hz for a total of 4 min). Total RNA was isolated using the AurumTM (Bio-Rad) RNA isolation kit according to the manufacturer's directions. For cellular samples, total RNA from NRVM was isolated using the high pure RNA isolation kit (Roche Applied Science) according to the manufacturer's directions. For qPCR, total RNA was deoxyribonuclease-treated on-column and transcribed to complementary DNA using iScriptTM (Bio-Rad) following the manufacturer's protocol. qPCR was performed with the TaqMan method (using the Roche Universal Probe Library System) on an ABI Step One Plus real-time PCR system. Relative expression was calculated using the $\Delta\Delta Ct$ method with normalization to constitutive genes as indicated in each figure. Specific primer/probe sequences are available on request.

Co-Immunoprecipitation-Co-IP was performed from nuclear protein extracts. Nuclear protein from 293 cells or mouse tissues was prepared using the Ne-PER kit (Pierce) according to the manufacturer's instructions. For each IP, 100-200 μ g of nuclear protein was loaded in IP dilution buffer (50 mм Tris-HCl, pH 7.4, 150 mм NaCl, 0.2% Nonidet P-40, supplemented with protease inhibitors) and immunoprecipitated with $2-5 \mu g$ of antibody bound to Dynabeads. Antibodies used for IP were anti-Myc (9E1, Santa Cruz Biotechnology) and goat polyclonal anti-KLF15 (Abcam). Immune complexes were washed extensively and eluted in SDS sample buffer. Immunoprecipitated and input proteins were run on SDS-PAGE and immunoblotted with the following antibodies as indicated: anti-p300 (N-15, Santa Cruz Biotechnology), anti-HA (HA-7, Sigma), anti-Myc (9E10, Santa Cruz Biotechnology), and goat polyclonal anti-KLF15 (Abcam). Secondary HRP-conjugated antibodies and ECL-plus chemiluminescent detection reagent were from Amersham Biosciences.

Microarray Analysis and Bioinformatics-Total ventricular tissue RNA was purified from wild-type and $Klf15^{-/-}$ mice (n = 4), which were subject to an 18-h overnight fast (4 p.m. to 10 a.m. the following morning). Labeling, hybridization, and scanning using an Affymetrix Genechip (Mouse Genome 430A 2.0 Array) were performed at the Harvard Partners Center for Genetics and Genomics. Raw microarray data were subject to background subtraction and multiarray normalization. Microarray data were log-transformed before differential expression analysis using the limma package (37). Probes with a p value of less than 0.05 after adjusting for false discovery rate using Benjamini and Hochberg method (38) were considered significant in differential expression. Enrichment in biological processes was analyzed using the DAVID bioinformatics suite (39, 40). The data set was submitted to the NCBI Gene Expression Omnibus with the accession number GSE53231.

Promoter-Luciferase Studies—Transient transfections of the indicated promoter plasmids using cultured NRVM or C2C12 cells at 70% confluence were carried out using Fugene6 (Roche



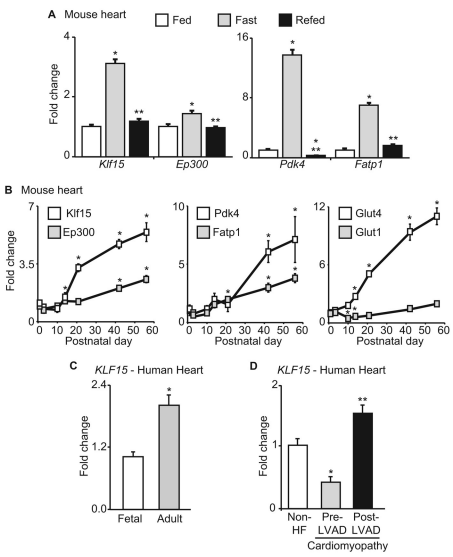


FIGURE 1. **KLF15 is regulated in physiological and pathological conditions.** *A*, *Klf15*, *Ep300*, *Pdk4*, and *Fatp1* expression in mouse heart during fed, fasted (24 h), and refed (24 h of fasting followed by 24 h of refeeding). n = 5. *, p < 0.05 versus Fed. **, p < 0.05 versus fasted. *B*, *Klf15*, *Ep300*, *Pdk4*, *Fatp1*, *Glut4*, and *Glut1* expression in mouse heart during postnatal maturation (n = 4). *, p < 0.05 versus day 1. *C*, *KLF15* expression in fetal versus adult human heart tissue (n = 3-4). p < 0.05. *D*, *KLF15* expression in human hearts that are non-failing, with advanced heart failure (prior to LVAD placement), and post-LVAD placement (n = 3-4). *, p < 0.05 versus non-HF. **, p < 0.05 versus Pre-LVAD. Rodent values are normalized to cyclophilin-B (*Ppib*) and human values to 18 S rRNA. *Error bars*, S.E.

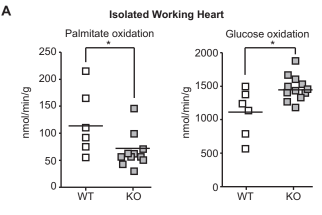
Applied Science and Promega) following the manufacturer's instructions, and bioluminescence was recorded on a Veritas luminometer (Turner Biosystems) as described previously (20, 26).

Data Analysis—All results are expressed as means, and *error* bars depict S.E. Statistical analysis of microarray data is described separately above. For experiments comparing the means of two normally distributed groups, two-tailed Student's t test for unpaired data was used. Statistical significance was defined as p < 0.05.

RESULTS

KLF15 Regulates Myocardial Lipid Utilization—Because KLF15 has been demonstrated to be important for fasting-induced gluconeogenesis in the liver (24), we first assessed its expression pattern in hearts of adult mice in the fed, fasted, and refed states. As with previously identified metabolic regulators, such as *Pdk4* and *Fatp1*, *Klf15* was robustly induced with fasting with a return to baseline after refeeding (Fig. 1A), suggesting a role in cardiac metabolic function. Klf15 expression was very low in neonatal mouse hearts and significantly induced with postnatal maturation in a pattern that paralleled other important metabolic genes (Pdk4 and Fatp1) during this transition from neonatal cardiac metabolism (glucose utilization) to that of the adult (predominance of lipid oxidation) (7) (Fig. 1B). In a similar manner, KLF15 expression was induced during the fetal to adult transition in human heart tissue (Fig. 1C). Conversely, KLF15 has been shown to be substantially down-regulated in rodent models of heart failure (20, 21), a pathologic state where myocardial lipid utilization is reduced (5, 6). Consistent with this rodent data, we found KLF15 to be significantly reduced in failing human hearts with a significant recovery of expression after mechanical unloading with a left ventricular assist device (Fig. 1D) (6). Although we have shown that KLF15 functions as a transcriptional repressor of prohypertrophic signaling in cardiomyocytes (20, 21), the gene networks directly regulated by





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Parameter	WT (n=8)	KO (n=12)	p-value
Heart rate (min-1)	242 <u>+</u> 14	263 <u>+</u> 9	0.20
Peak systolic pressure (mm Hg)	69 <u>+</u> 2	71 <u>+</u> 1	0.59
Cardiac Output (mL/min)	10.4 <u>+</u> 1.7	13.9 <u>+</u> 1.2	0.11
Aortic Output (mL/min)	8.2 <u>+</u> 1.3	10.7 <u>+</u> 0.9	0.12
Coronary flow (mL/min)	2.9 <u>+</u> 0.4	3.1 <u>+</u> 0.3	0.70
Cardiac work (mm Hg x mL/min)	7.1 <u>+</u> 1.3	9.7 <u>+</u> 0.8	0.10

FIGURE 2. **KLF15 is essential for myocardial lipid utilization.** *A*, fatty acid (palmitate) and glucose oxidation rates in WT versus KO hearts in isolated working heart preparation (n = 8 - 12). *, p < 0.05 versus WT. *B*, hemodynamic parameters in isolated working heart studies. WT and KO hearts (n = 8 - 12) were perfused in isolated working mode, and hemodynamic parameters were measured as listed. There was no statistically significant difference in hemodynamic performance between genotypes.

KLF15 in the myocardium remain largely unknown. Based on the data above, we hypothesized that KLF15 might play an important role in metabolic function of the adult heart.

To establish a tissue-intrinsic role for KLF15 in cardiac metabolism, we performed ex vivo isolated working heart (IWH) studies in wild-type (WT) and *Klf15*-deficient (KO) hearts. We focused these IWH studies on fatty acid and glucose utilization because these substrates are the major fuel sources in the working mammalian heart (1). KO hearts had a striking reduction in fatty acid oxidation with a parallel increase in glucose oxidation (Fig. 2A). Importantly, there was no significant difference in hemodynamic indices between KO and WT hearts during the IWH experiments (Fig. 2B). Furthermore, KO hearts are structurally normal and have preserved systolic function under basal conditions (20). These results suggest that the observed defect in lipid utilization is not occurring during a state of severe contractile dysfunction. In addition, KO mice are characterized by reduced intramyocardial FA and TG abundance (Fig. 3A), despite elevated plasma concentration of FAs and TGs (Fig. 3B). This observation was corroborated by electron microscopic examination, which revealed a striking paucity of intramyocellular lipid droplets in the hearts of KO mice (Fig. 3C). Hence, KLF15 deficient hearts are unable to ade-

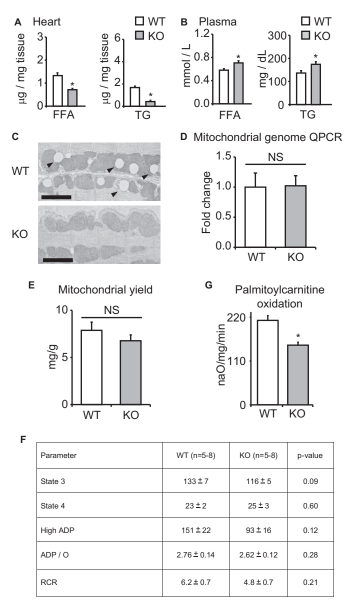


FIGURE 3. **Mitochondrial parameters in KLF15 KO.** Shown are FA and TG content in WT *versus* KO heart tissue (A) and plasma (B) (n = 5). *, p < 0.05 *versus* WT. C, representative transmission electron micrographs of WT *versus* KO heart tissue demonstrating marked paucity of intramyocellular lipid droplets (*arrowheads*) in KO tissue. *Bar*, 2 μ m. *D*, qPCR for relative mitochondrial genome content in WT *versus* KO heart tissue (n = 4). Values are normalized to *Rplp0* (*36B4*) genomic DNA. *E*, mitochondrial yield from whole heart tissue of WT *versus* KO mice, expressed as mg of mitochondrial protein/g of heart tissue (n = 4). *F*, rates of glutamate oxidation in mitochondria freshly isolated from WT *versus* KO hearts (n = 5-8). *G*, palmitoylcarnitine oxidation rates in freshly isolated mitochondria from WT *versus* KO hearts (n = 4). *, p < 0.05. *Error bars*, S.E.

quately partition plasma lipids despite adequate delivery to the myocardium.

To demonstrate an additional downstream defect in mitochondrial lipid catabolism independent of the above defect in cytosolic lipid bioavailability, we directly assessed substrate utilization in mitochondria freshly isolated from the hearts of WT and KO mice. We did not detect differences in mitochondrial genome copy number in KO hearts (Fig. 3*D*). Concordant with this finding, mitochondrial yield was equivalent between WT and KO hearts, thus providing additional confirmation that



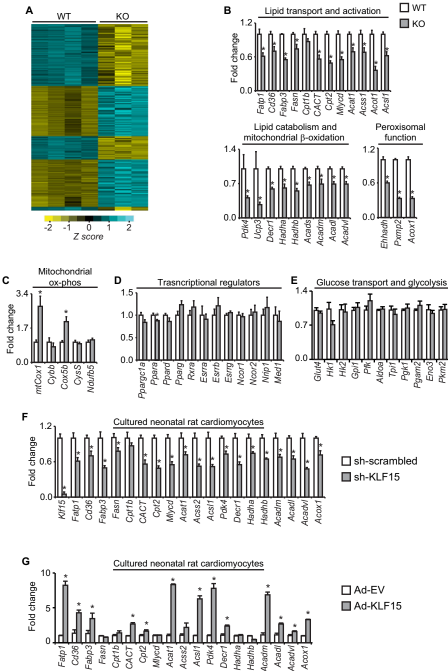


FIGURE 4. **KLF15 regulates the myocardial lipid flux gene program.** *A*, microarray heat map of transcript expression profiles from WT/KO hearts (n = 3-4). *B*, expression (qPCR) in WT *versus* KO hearts of critical transcripts involved in myocardial lipid flux; *C*, mitochondrial oxidative phosphorylation; *D*, transcriptional regulation; *E*, glycolysis (n = 4-5). *, p < 0.05. Expression (qPCR) of a parallel panel of lipid flux transcripts in cultured neonatal rat ventricular cardiomyocytes after shRNA-mediated *Klf15* silencing (*F*) and adenovirus-mediated *Klf15* overexpression (*G*) (n = 5-6). *, p < 0.05. Values normalized to *Ppib. Error bars*, S.E.

mitochondrial quantity was unchanged (Fig. 3*E*). Furthermore, WT and KO cardiac mitochondria exhibited no significant differences in glutamate oxidation rates, reflecting unaltered function of the electron transport chain (Fig. 3*F*). Last, transmission electron microscopy at the mitochondrial level revealed no gross differences in mitochondrial ultrastructure (Fig. 3*C*). However, when we directly assessed mitochondrial lipid oxidation rates, we found that freshly isolated mitochondria from KO hearts had a significant defect in fatty acid oxidation (Fig. 3*G*). Thus, in addition to proximal abnormalities in sarcolemmal

lipid partitioning (Fig. 3, *A* and *B*), KLF15 is also critical for the distal step of mitochondrial lipid oxidation.

To gain further insight into the broad defects in myocardial lipid utilization seen above, we performed transcriptome-wide gene expression profiling of KO heart tissue. These microarray analyses showed that KO hearts have a transcriptomic signature characterized by a broad and significant reduction in genes critical for myocardial substrate metabolism, including genes involving lipid utilization (Fig. 4*A* and supplemental Table 1). Because lipids are the dominant fuel source for the heart under



physiologic conditions, the above transcriptomic signature and our IWH data (Fig. 2A) prompted us to directly assess a broad panel of lipid-related transcripts by qRT-PCR in WT versus KO heart tissue. These analyses confirmed significant reductions in genes encompassing nearly all major components of the myocardial lipid flux pathway, including sarcolemmal FA partitioning/activation, intramyocellular storage, mitochondrial transport and β -oxidation, and peroxisomal function (Fig. 4B). Genes directly involved in the mitochondrial electron transport chain and oxidative phosphorylation were not reduced in KO hearts (Fig. 4C). Importantly, the observed reductions in lipid flux targets seen in KO hearts occurred without significant alterations in the expression of known transcriptional regulators of myocellular substrate flux and energetics (Fig. 4D). Moreover, we did not observe significant differences in key genes involved in myocardial glucose uptake and glycolysis (Fig. 4E) (12, 16, 41). Acute silencing as well as overexpression of KLF15 in cultured neonatal rat cardiomyocytes significantly reduced and augmented, respectively, expression of a parallel panel of lipid flux targets, confirming a cell-autonomous role for KLF15 in cardiomyocyte gene expression (Fig. 4, G and H).

KLF15 Transactivates Direct Targets via Recruitment of p300— We next sought to establish the principle that KLF15 could regulate key targets in this pathway via a direct mechanism of action. *Pdk4* and *Fatp1* (*Slc27a1*) were used as exemplary targets, given the essential role of these genes in myocellular lipid flux and extensive functional knowledge of their proximal promoters (30, 42–45). The proximal promoters of both genes were induced by KLF15 in cultured cardiomyocytes (Fig. 5*A*). ChIP analysis in cardiomyocytes confirmed enrichment of KLF15 at conserved KLF response elements on these promoters (Fig. 5*B*). Thus, although *Fatp1* and *Pdk4* are only representative of a larger array of KLF15-regulated metabolic targets, these focused analyses serve to establish the principle that KLF15 can regulate important genes via direct occupancy of target promoters.

In order to gain deeper insight into a generalizable mechanism underlying KLF15-mediated target gene transactivation, we turned to structure-function analyses. We were intrigued by a recent biophysical study that demonstrated significant similarity between the p300-interacting domain of p53 and a homologous domain within the erythroid-specific KLF family member, EKLF/KLF1 (46). Indeed, inspection of the KLF15 gene revealed a putative p300-interacting transactivation domain (Fig. 5C) that was highly conserved from humans to zebrafish (Fig. 5D). The possibility of cooperativity between these two regulators on metabolic targets was further supported by our observation that myocardial Ep300 and Klf15 expression were induced in parallel during fasting and postnatal maturation (Fig. 1, A and B). Indeed, we observed significant cooperativity between KLF15 and p300 upon induction of target promoters (Fig. 5*E*). Studies in $p300^{-7}$ cells revealed that optimal target gene induction by KLF15 required endogenous p300 (Fig. 5F). We next demonstrated interaction between epitope-tagged KLF15 and p300 when co-expressed in 293 cells and also confirmed interaction between endogenous KLF15 and p300 in mouse heart and liver tissues (Fig. 6, A-C). Ablation of the KLF15 putative p300 binding domain (KLF15 Δ p300)

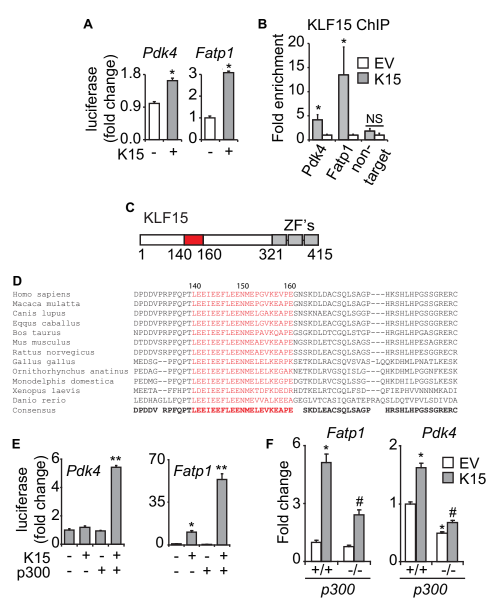
abrogated the interaction with endogenous p300 (Fig. 6*C*) and the cooperative induction of target promoters (Fig. 6*D*). Importantly, KLF15 Δ p300 remains nucleus-localized and is expressed as a protein of expected mobility with abundance identical to that of identical full-length KLF15 (Fig. 6*C*). Finally, ChIP against endogenous p300 in KLF15-deficient mouse heart tissue revealed reduced p300 enrichment in the proximal promoters of target genes (Fig. 6*E*). Taken together, the above data establish KLF15 as a direct regulator of the myocardial lipid flux pathway *in vivo*. In addition, we demonstrate that a major mechanism of KLF15-mediated transactivation involves interaction with p300 and site-specific recruitment of this critical co-activator to target promoters.

DISCUSSION

Our current findings establish KLF15 as a direct and independent regulator of myocardial nutrient flux and thereby introduce an entirely new class of factors (KLFs) as core components of the transcriptional circuitry that governs cardiac metabolism (schematized in Fig. 7). Using unbiased gene expression profiling, we elucidate a large number of KLF15-dependent targets in the heart in vivo and confirm its essential role in regulating a transcriptional program essential for efficient myocardial lipid flux. Furthermore, we establish a fundamental molecular mechanism of KLF15-mediated target gene transactivation, which involves its ability to interact with and recruit the chromatin remodeling enzyme p300. Because the mammalian heart's ability to maintain blood flow across a wide range of physiologic and pathophysiologic circumstances depends on tightly coupled control of fuel supply with energy demand (5), this work suggests that KLF15 is essential for the heart's ability to metabolically adapt to stress. Consistent with this critical role in myocardial adaptation, we have found that Klf15-deficient mice develop dramatically accelerated heart failure after hemodynamically (pressure overload) and neurohormonally (chronic angiotensin II infusion) induced stress (20, 21). When considered alongside our prior observations that implicate KLF15 as a regulator of fasting hepatic gluconeogenesis and skeletal muscle substrate catabolism (24-26), the current work implicates KLF15 as a nodal integrator of nutrient flux across diverse organ systems.

We show that cardiac KLF15, which is robustly induced during physiologic states, such as fasting and postnatal development, cooperates with p300 to regulate key metabolic targets. Previous studies have implicated p300 acetyltransferase as a positive regulator of pathologic cardiomyocyte hypertrophy, possibly via its ability to acetylate prohypertrophic transcription factors, such as GATA4, MEF2, and p53 (21, 47-50). Prior work from our group has shown that Klf15 expression is dramatically reduced during pathologic cardiac hypertrophy, an event that releases repression of GATA4, MEF2, and the p300p53 transcriptional module (20, 21). Therefore, our current findings, when taken in the context of previously published work, support a model in which increased levels of KLF15 (e.g. during fasting) favor recruitment of p300 to specific metabolic targets, whereas reduced levels of KLF15 (e.g. during pressure overload) decrease expression of these metabolic genes and simultaneously liberate molecules of p300 to co-activate prohy-





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FIGURE 5. **KLF15 directly regulates metabolic targets.** *A*, induction of *Pdk4*(-1.6kb)-luc and *Fatp1*(-1.2kb)-luc in transfected NRVM (n = 6). *, p < 0.05. *B*, ChIP against exogenously expressed KLF15 in NRVM on the proximal promoter regions of the endogenous rat *Pdk4* and *Fatp1* loci in the vicinity of highly conserved KLF consensus sites (n = 3). *, p < 0.05. *C*, domain map of KLF15 protein depicting putative p300-interacting transactivation domain (residues 140–160). *ZF*, zinc finger. *D*, alignment of KLF15 protein sequence across species in the vicinity of highly conserved p300-interacting transactivation domain (residues 140–160). *E*, cooperative promoter induction between KLF15 and p300 on *Pdk4*(-1.6kb)-luc and *Fatp1*(-1.2kb)-luc in C2C12 cells (n = 5). *, p < 0.05 *versus* mock transfection. **, p < 0.05 versus K15 full-length. *F*, expression of the indicated transcripts in p300^{+/+} versus p300^{-/-} MEFs after KLF15 overexpression (n = 5). Values normalized to *Ppib*. *, p < 0.05 versus K15 p300^{+/+}. #, p < 0.05 versus K15 p300^{+/+}.

pertrophic transcription factors, such as GATA4 and MEF2. In this manner, increased levels of KLF15 may recruit rate-limiting quantities of p300 to transactivate its direct targets while simultaneously repressing the function of p300-regulated prohypertrophic transcriptional effectors.

Our IWH experiments show that *Klf15*-deficient hearts have not only a 47% reduction in palmitate oxidation but also a 32% increase in glucose oxidation rate. These IWH data parallel those observed in *Ppara* null hearts (15). Although our studies demonstrate that KLF15, like PPAR α , directly regulates genes critical for efficient myocardial lipid utilization, there are several potential explanations for the concomitant increase in glucose oxidation rates. We did not observe increased expression of key genes that directly participate in myocardial glucose uptake or glycolysis (Fig. 4*E*). However, KO hearts did have reduced expression of *Pdk4*, an important inhibitor of the pyruvate dehydrogenase complex. Decreased PDK4 would increase pyruvate dehydrogenase activity, leading to increased rates of glucose oxidation. In addition, it is possible that a primary reduction in myocardial long chain fatty acid oxidation rates can secondarily increase glucose oxidation rates via the well described allosteric mechanisms of the Randle cycle (51). Finally, it remains possible that KLF15 might regulate the metabolism of other myocardial fuels (*e.g.* amino acids) in a manner that could influence glucose homeostasis.

Because cardiac KLF15 is induced during fasting *in vivo* (Fig. 1*A*), delineation of the precise molecular events that govern its expression will provide additional insights into its physiologic



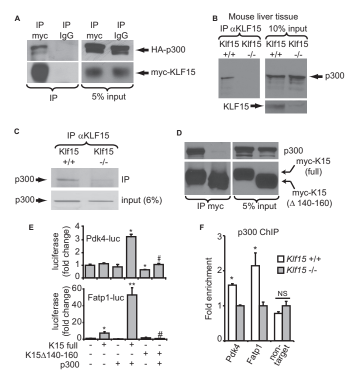


FIGURE 6. **KLF15 and p300 interact to regulate metabolic targets.** *A*, co-IP of heterologously expressed, epitope-tagged KLF15 and p300 in 293 cells. *B*, co-IP of endogenous KLF15 and p300 in mouse liver tissue nuclear extracts. *C*, co-IP of endogenous KLF15 and p300 in mouse heart tissue nuclear extracts. *Klf15^{-/-}* tissue was used as a negative control for co-IPs. *D*, co-IP between overexpressed KLF15 and endogenous p300 in nuclear extracts of 293 cells. *E*, transfections demonstrating loss of cooperativity with K15Δ140–160 in C2C12 cells. *n* = 5. *, *p* < 0.05 *versus* mock transfection. **, *p* < 0.05 *versus* K15 full-length. +, *p* < 0.05 *versus* K15 full-length. +, *p* < 0.05 *versus* K15 m the proximal promoters of *Pdk4* and *Fatp1* in the vicinity of the KLF15 binding site (*n* = 3). *, *p* < 0.05. *Error bars*, S.E.

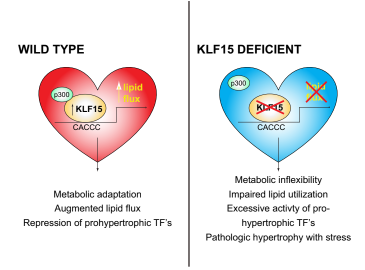


FIGURE 7. Schematic representing KLF15 as a regulator of cardiac lipid flux.

function. Recent studies by our group and others have identified several neurohormonal pathways as important regulators of KLF15 expression in non-cardiac tissue. Signals known to drive nutrient catabolism, such as glucocorticoids (52, 53) and cAMP-response element-binding protein activation (54), have been shown to induce KLF15 expression, whereas anabolic sig-

nals, such as insulin, can suppress its expression (55). In addition, we have recently demonstrated that cardiac KLF15 expression is directly regulated by BMAL1 (27), a central component of the mammalian circadian clock. Therefore, it is likely that balance between such catabolic and anabolic upstream signals, with additional synchronization via the circadian clock, regulates cardiac KLF15 function. In addition to its newly defined role in cardiac lipid metabolism, KLF15 also regulates the catabolism of skeletal muscle branched chain amino acids, the body's major store of glucogenic precursors (24, 25). Because cardiac and skeletal muscle share similar substrate flux machinery, it is possible that KLF15 might also regulate myocardial amino acid homeostasis in a manner that influences metabolism of other substrates and cardiac function. Given the intense interest in the interplay between lipid and branched chain amino acid metabolism during physiological and pathophysiological states (56), it will be interesting to explore whether these aspects of KLF15 function are perturbed in heart failure and other cardiometabolic diseases.

Our current work in cardiac metabolism is also interesting to consider alongside our recent observation that KLF15 regulates cardiac repolarization *in vivo* (27). Studies in large animal models of heart failure demonstrate that regional electromechanical dyssynchrony is associated with profound changes in genes critical for myocardial lipid and amino acid metabolism (57). Because KLF15 expression has also been shown to be significantly reduced in rodent and human heart failure (21) with recovery of expression after mechanical unloading (Fig. 1*D*), these findings collectively raise the intriguing possibility that myocardial substrate flux and electrical activity might be subject to coordinate control by KLF15 as a form of "metabo-electrical coupling."

Previous studies have identified key members of the nuclear receptor superfamily (e.g. PPARs and ERR α) as direct regulators of cardiac lipid metabolism (12, 15). We note that many of the lipid metabolic genes regulated by KLF15 are also known to be PPAR α targets in the heart (12). Furthermore, we find that KLF15-deficient hearts have defects in lipid utilization as prominent as those observed with complete PPAR α deficiency (15). Importantly, these defects in lipid metabolism and target gene expression seen with KLF15 deficiency occur with minimal alteration in expression of PPARs, ERRs, or several other important metabolic regulators (Fig. 4D). We postulate that KLF15 and nuclear receptors, such as PPAR α , together exert coordinated control over a subset of important shared targets. Our observation that KLF15 binds and recruits p300 to promoters supports a transactivation model in which KLF15 occupancy favors an open chromatin configuration for the regulatory regions of its direct targets in the lipid flux pathway. It is also possible that the ability of KLF15 to influence intracellular lipid abundance may secondarily affect the bioavailability of endogenous PPAR ligands (13). Studies exploring potential cross-talk between KLFs and nuclear receptors in cardiac metabolism are an area of ongoing investigation by our group. Interestingly, KLF homologs and PPAR α -like nuclear receptor genes in C. elegans have recently been shown to both play important roles in lipid metabolism (58, 59), raising the intriguing possibility that coordinated control of metabolic flux by



these two ancient transcription factor families may have been present as early in evolution as the phylum *Nematoda*. We speculate that functional interactions between KLFs and nuclear receptor networks will emerge as an important theme in our understanding of metabolic plasticity in the myocardium as well as across a broad range of tissues.

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