Octanoate is differentially metabolized in liver and muscle and fails to rescue cardiomyopathy in CPT2 deficiency

Andrea S. Pereyra¹, Kate L. Harris², Arvin H. Soepriatna³, Quin A. Waterbury⁴, Sivakama S. Bharathi⁴, Yuxun Zhang⁵, Kelsey H. Fisher-Wellman¹, Craig J. Goergen⁶, Eric S. Goetzman⁷, and Jessica M. Ellis¹⁸

¹Brody School of Medicine at East Carolina University, Department of Physiology, and East Carolina Diabetes and Obesity Institute, Greenville, NC, USA; ²Department of Biochemistry and ³Weldon School of Biomedical Engineering, Purdue University, West Lafayette, IN, USA; and ⁴Department of Pediatrics, Children's Hospital of Pittsburgh of the University of Pittsburgh Medical Center, University of Pittsburgh School of Medicine, Pittsburgh, PA, USA

Abstract

Long-chain fatty acid oxidation is frequently impaired in primary and systemic metabolic diseases affecting the heart; thus, therapeutically increasing reliance on normally minor energetic substrates, such as ketones and medium-chain fatty acids, could benefit cardiac health. However, the molecular fundamentals of this therapy are not fully known. Here, we explored the ability of octanoate, an eight-carbon medium-chain fatty acid known as an unregulated mitochondrial energetic substrate, to ameliorate cardiac hypertrophy in long-chain fatty acid oxidation-deficient hearts because of carnitine palmitoyltransferase 2 deletion (Cpt2⁻/⁻). CPT2 converts acyl-carnitines to acyl-CoAs in the mitochondrial matrix for oxidative bioenergetic metabolism. In Cpt2⁻/⁻ mice, high octanoate-ketogenic diet failed to alleviate myocardial hypertrophy, dysfunction, and acylcarnitine accumulation suggesting that this alternative substrate is not sufficiently compensatory for energy provision. Aligning this outcome, we identified a major metabolic distinction between muscles and liver, wherein heart and skeletal muscle mitochondria were unable to oxidize free octanoate, but liver was able to oxidize free octanoate. Liver mitochondria, but not heart or muscle, highly expressed medium-chain acyl-CoA synthetases, potentially enabling octanoate activation for oxidation and circumventing acylcarnitine shuttling. Conversely, octanoylcarnitine was oxidized by liver, skeletal muscle, and heart, with rates in heart 4-fold greater than liver and, in muscles, was not dependent upon CPT2. Together, these data suggest that dietary octanoate cannot rescue CPT2-deficient cardiac disease. These data also suggest the existence of tissue-specific mechanisms for octanoate oxidative metabolism, with liver being independent of free carnitine availability, whereas cardiac and skeletal muscles depend on carnitine but not on CPT2.

Because the heart is a continuously working muscle, it has a high bioenergetic demand and relies primarily on fatty acids as substrate. Fatty acids are particularly important cardiac substrates as indicated by dysregulated mitochondrial fatty acid oxidation (mFAO) in metabolic diseases, such as diabetes, obesity, hypertension, and cardiovascular disease. As such, cardiac energy insufficiency continues to be recognized as the underlying etiology of clinical hypertrophic cardiomyopathy (1). Cardiac hypertrophic remodeling is an independent predictor of all-cause death (2, 3) and a common occurrence in obese (4, 5), hypertensive (6), diabetic (7), and aged (8) individuals, and it is well established that it occurs as a direct consequence of impaired oxidative metabolism in the myocardium. In agreement, animal models of cardiac hypertrophy have documented energetic deficits and/or disruptions in mFAO (9–11). Moreover, cardiac hypertrophy occurs as a direct consequence of primary defects in mFAO (12–16). The metabolic pathway of beta-oxidation of long-chain fatty acids for energy is regulated by the transport of fatty acids into the mitochondrial matrix via the carnitine shuttle. The first and required step in mitochondrial long-chain fatty acid oxidation is the activation by acyl-CoA synthetase, being ACSL1 the predominant activator for oxidative metabolism in the heart (13, 17–20). Once the acyl-CoA moiety is formed, the CoA group is exchanged for a carnitine molecule by the rate-limiting enzyme carnitine palmitoyltransferase 1 (CPT1) to generate an acylcarnitine. The acylcarnitine is then transported through the inner mitochondrial membrane by the carnitine-acylcarnitine translocase (CACT). Once the acylcarnitine is inside the mitochondria, carnitine palmitoyltransferase 2 (CPT2) exchanges the carnitine group for a mitochondrial CoA to recreate an acyl-CoA for subsequent beta-oxidation.

In addition to mFAO, the high-energy demands of the heart are also met by glucose oxidation. Indeed, the loss of cardiac glucose oxidation because of deletion of...
the mitochondria pyruvate carrier subunits results in cardiac hypertrophy (21, 22). Remarkably, this severe cardiac hypertrophy is rescurable by high-fat or ketogenic medium-chain fatty acid diets (21, 22) suggesting that ketones, long-chain fatty acids, and medium-chain fatty acids could serve as critical alternative substrates for cardiac oxidative metabolism. We and others have demonstrated that loss of myocardial mFAO by the genetic deletion of CPT2 or CPT1 in mouse myocardium results in cardiac remodeling, hypertrophy, and heart failure (14, 23). In patients with severe genetic defects of Cpt2, cardiomyopathy is common, and medium-chain fatty acid-enriched diet is used as therapy. Medium-chain fatty acids of eight carbons or fewer provide an alternative energetic substrate because they are thought to bypass the carnitine-dependent CPT1-CACT-CPT2 shuttling system to access the mitochondrial matrix for subsequent oxidation. Here, we tested the ability of the medium-chain fatty acid octanoate to improve CPT2 deficiency-induced cardiac hypertrophy. Surprisingly, a heavily octanoate-enriched ketogenic diet was not able to alleviate CPT2 deficiency-mediated cardiac hypertrophy. This effect is potentially mediated by limited access of the heart to dietary octanoate, as the liver receives the first pass of exogenous octanoate through the portal vein upon intestinal absorption. Here, we found that liver has an additional metabolic advantage because liver mitochondria were able to oxidize free octanoate, whereas the heart and skeletal muscle were not. On the contrary, the carnitine-bound form of octanoate was readily oxidized by heart, skeletal muscle, and liver mitochondria evidencing that the specific octanoate metabolites are oxidized distinctly in a tissue-dependent manner. Despite the requirement of carnitine for octanoate oxidation in muscles, octanoylcarnitine oxidation rates are not impacted by CPT2 deficiency. These data suggest that free octanoate can be directly oxidized in liver but not in cardiac and skeletal muscles. Furthermore, in muscles, octanoate oxidation is carnitine dependent but CPT2 independent. Our data also suggest that mFAO is obligatory for maintaining cardiac size and function.

MATERIALS AND METHODS

Animal

Heart- and muscle- (Cpt2<sup>−/−</sup>) and skeletal muscle-specific (Cpt2<sup>−/−</sup>/mCK-cre) CPT2 conditionally deficient mice were generated using Cre recombinase driven by the muscle creatine kinase promoter (Jackson Laboratories; stock no. 006475) or the human alpha-skeletal actin promoter (The Jackson Laboratory; stock no. 006149) as described (23, 24). Littermates lacking the Cre transgene were used as controls. Mice were given free access to water and standard chow (PicoLab 5053; Lab Diets), in pathogen-free housing under 12-h light-dark cycles. All procedures were approved by the Purdue Animal Care and Use Committee (assurance A3231-01) and the Institutional Animal Care and Use Committee of East Carolina University (assurance A3469-01). For diet studies, mice were given ad libitum access to either control (PicoLab 5053; Lab Diets, Richmond, IN; TD94045; Envigo Teklad, Madison, WI), ketogenic diet (F6689; fat:protein + carbohydrate, 43; BioServ, Flemington, NJ), or trioctanoin-enriched diet (TD70585; Envigo Teklad). Trioctanoin (Captex 8000) obtained from Abitec was used for customized diets synthesized by Envigo (supplemental Table S1). All dietary interventions began between week 3 and 6 of age and lasted 4 weeks. For carnitine studies, powdered i-carnitine (Sigma C9283) was dissolved in the drinking water at 300 mg/kg/day, as described (24, 25). Echocardiograms were collected as previously described (23) from isoflurane-anesthetized mice (1–3% in 1.5 L/min medical-grade air) via high-frequency ultrasound (Vevo2100; FUJIFILM VisualSonics). M-mode, B-mode, and respiratory- and cardiac-gated images (electrocardiographically gated kilohertz visualization) were collected for both medial short- and long-axis views (MS550D transducer; FUJIFILM VisualSonics). Echocardiogram-derived calculations were determined from the Endocardial and Epicardial Area Protocol (Vevo Lab; FUJIFILM VisualSonics). Beta-hydroxybutyrate was measured using a commercially available kit according to manufacturer’s instruction (StanBio).

Lipid analysis

Lipid profiling of tissue and biological fluids was performed as previously described (23). Briefly, lipids were extracted from tissues using Bligh and Dyer method (26), and both the lipid and polar phases were dried separately, resuspended, and injected directly via a microautosampler (G1377A) into a QQQ6410 triple quadrupole mass spectrometer (Agilent Technologies, San Jose, CA) operated in the positive-ion mode and equipped with ESI source ion source, as have described (23, 27). Data were collected in multiple reaction monitoring mode (23, 27, 28), and ion intensities acquired by an in-house script were further normalized to sample protein concentration.

Molecular response and indicators

RNA was isolated using Trizol (Invitrogen), and RNA was converted to complementary DNA (Applied Biosystems High Capacity cDNA RT Kit) and used for SYBR Green (Bio-Rad)-based real-time PCR. Results were normalized to housekeeping gene and expressed as arbitrary units of $2^{−ΔΔCT}$. For Western blots, lysates were collected in lysis buffer (50 mM Tris-HCl, 150 mM NaCl, 1 mM EDTA, and 1% Triton X-100) with protease and phosphatase inhibitors. Homogenates, lysates, or mitochondrial pellets were equally loaded and electrophoresed on SDS-polyacrylamide gels, transferred to nitrocellulose membrane, blocked with 5% milk-Tris-buffered saline with 0.1% Tween® 20 detergent for 1 h, incubated with primary antibody (1:1000–12,000) against alpha-tubulin (Sigma T0198; pyruvate dehydrogenase (PDH) and phospho-PDH (S293) (Cell Signaling 3205 and 31866); medium-chain acyl-CoA synthetase 3 (ACSM3; Invitrogen PA5-100374); alpha-AMP-activated protein kinase and phospho-AMP-activated protein kinase (T172) (Cell Signaling 5831 and 2535), washed, and incubated with secondary antibodies conjugated to infrared dye 800CW or 680LT (LiCor). Proteins were visualized with Odyssey and quantified using Image Studio (LiCor). Carnitine O-octanoyltransferase (cOT) expression in isolated mitochondria from mouse liver and heart was determined by label-free proteomics via nano LC-MS/MS as described previously (29).
Oxidation assays
Oroboros high-resolution respirometry was performed on fresh whole-tissue homogenate and/or isolated mitochondria from wild-type C57Bl/6j mouse liver, heart, and skeletal muscle and Cpt2M−/− skeletal muscle. Homogenization was performed in Mir05 buffer (MgCl2·6H2O 5 mM, K⁺MES 105 mM, taurine 20 mM, KH2PO4 10 mM, Hepes 20 mM, d-sucrose 110 mM, and fatty acid-free BSA 1 g/l) at 20 or 40 w/v. Mitochondria were isolated by standard differential centrifugation, and yield was assessed by measuring protein concentration in mitochondrial pellets (30). Respiration was measured in Mir05 media (31) using 60 μl of tissue homogenate, 100 μg (liver) or 50 μg (heart and skeletal muscle) of isolated mitochondria in each chamber. After baseline respiration was established, malate (2 mM; Sigma M1296) was added to maintain the tricarboxylic acid cycle, followed by ADP (2 mM) to stimulate respiration. Substrates were added in the following concentrations: free octanoate 0.2 mM (Sigma C5038), octanoyl-CoA 0.2 mM (Sigma O6877), octanoylcarnitine 0.2 mM (Sigma 50892), palmitic acid 0.02 mM (Sigma P0500), palmitoyl-CoA 0.02 mM (Sigma P9716), palmitoylcarnitine 0.02 mM (Sigma P61654), free CoA 0.125 mM (Sigma C5019), and l-carnitine HCl 5 mM (Sigma C0283). Complementary assays were performed in mitochondria isolated in KCl 100 mM, Mops 50 mM, EGTA 1 mM, MgSO4 5 mM, BSA 0.2%, buffer, pH 7.1, and respiration measured in ATP containing buffer (K-MES 105 mM, KCl 30 mM, KH2PO4 10 mM, MgCl2 5 mM, EGTA 1 mM, BSA 2.5 g/l, and pH 7.1). The experiments were repeated at least three times with mitochondria isolated from separate animals. To combine data from the three experiments, we calculated the ratio of ADP-stimulated fatty acid-induced respiration rates (lipid + malate in the presence of ADP) to that with malate alone (malate + ADP).

Medium-chain acyl-CoA dehydrogenase enzyme activity assays
Heart and liver lysates from C57Bl/6 mice were prepared in phosphate-buffered saline supplemented with 0.1% lubrol detergent. Medium-chain acyl-CoA dehydrogenase (MCAD) activity was determined using the electron-transferring flavoprotein fluorescence-based microplate assay exactly as described (32), using 25 μM octanoyl-CoA as substrate.

Statistics
Data are presented as mean ± SEM, unless otherwise specified. Statistical analysis and figures were generated using Excel or GraphPad Prism, version 8.00 for Windows (GraphPad Software). Data were compared using unpaired Student’s t-test and 1-way or 2-way ANOVA followed by multiple comparison analysis. Significance level was set at P < 0.05. Statistical details for each data set are described on figure legends. Heatmapper online tool was used to generate heat maps (Wishart Research Group).

RESULTS
Dietary octanoate does not attenuate cardiac hypertrophy induced by CPT2 deficiency
Individuals with inborn errors in mFAO, such as CPT2 deficiency, commonly supplement their diets with medium-chain fatty acids as an alternative fuel source. To determine if supplementation with octanoate would alleviate cardiac hypertrophy in CPT2 deficiency (Fig. 1A) by bypassing the carnitine-mediated transport shuttle for mitochondrial oxidation, we fed control and myocardial CPT2-deficient mice (Cpt2M−/−), a diet supplemented with trioctanoin, a triacylglycerol containing three 8-carbon-long acyl chains, as 20% wet weight in the regular diet. However, the octanoate-supplemented diet failed to rescue cardiac hypertrophy (Fig. 1B). Octanoate boluses and medium-chain ketogenic diets can increase circulating ketone bodies (33) because of stimulation of high rates of medium-chain mFAO flux in the liver. Importantly, medium-chain ketogenic diets can completely revert severe cardiac hypertrophy in models of impaired cardiac pyruvate oxidation (21, 34). Thus, to test the ability of highly enriched medium-chain ketogenic feeding to alleviate the cardiac phenotype in the absence of carnitine mediated-fatty acid oxidation, a diet was formulated with octanoate (OctD) as a majority of the diet. Octanoate, in the form of trioctanoin, comprised ~70% of the total fat and provided ~74% of the total dietary kilocalories (Fig. 1C). Control and Cpt2M−/− male and female mice were placed on either OctD or matched control diet (CD) at 3 weeks of age when neither cardiac hypertrophy nor heart failure is detectable in Cpt2M−/− mice (23). Dietary intervention lasted a total of 4 weeks. While Cpt2M−/− mice are mildly ketogenic on chow diet, as previously reported (23), the octanoate diet induced ketosis in both control and Cpt2M−/− mice (Fig. 1D). The octanoate diet neither altered body weight, by genotype, nor relative to the CD-fed mice (Fig. 1E) but did reduce adipose depot mass in both genotypes (Fig. 1F). Although not reaching significance, we observed a trend toward slightly reduced heart mass in the OctD-fed Cpt2M−/− mice compared with CD-fed Cpt2M−/− mice (Fig. 1G). Echocardiographic analysis confirmed that cardiac hypertrophy in Cpt2M−/− mice was not attenuated by OctD as indicated by sustained elevated left ventricular mass (Fig. 2A).

We previously reported that Cpt2M−/− mice had compromised cardiac function (23). Here, Cpt2M−/− mice fed either chow or OctD presented with increased end-diastolic volume along with a 25% reduction in flow velocity as indicated by Doppler analysis (Fig. 2B, C). Ejection fraction remained significantly lower in Cpt2M−/− mice independent of diet (Fig. 2C). In Cpt2M−/− mice, the OctD slightly increased peak-systolic volume, whereas in control mice, the OctD increased end-diastolic volume, stroke volume, and peak-systolic volume by ~50% (Fig. 2D). Interestingly, while cardiac output was not changed in Cpt2M−/− mice with OctD, control mice had a ~1.5-fold increased output (Fig. 2D). Together, these data demonstrate that the octanoate-rich diet modulates cardiac function of control mice, an effect that Cpt2M−/− mice are impervious to, and that octanoate diet fails to alleviate cardiac hypertrophy or to improve cardiac function in CPT2-deficient hearts.
Alternative dietary fuels regulate cardiac hypertrophy-related genes

Next, we assessed the molecular response of Cpt2M−/− hearts to octanoate dietary intervention. OctD modestly reduced cardiac expression of atrial natriuretic factor and increased actin alpha 1 expression in Cpt2M−/− mice, thus inducing minimal changes on molecular indicators of pathological remodeling compared with CD-fed Cpt2M−/− mice (Fig. 3A). OctD did not trigger expression of hypertrophy genes in control mice (Fig. 3A, B). The mechanistic target of rapamycin (mTOR) growth pathway is a critical regulator of cardiac hypertrophy (13, 35, 36). While Cpt2M−/− hearts had increased mTor, reduced autophagy marker Atg10, and increased phosphorylation of downstream target p70S6K of mTOR, the OctD did not significantly affect these measures for either genotype (Fig. 3B and supplemental Fig. S1A). We previously showed that a
In relation to mitochondrial crosstalk with other tissues and organs, myomitoines are predicted to regulate systemic metabolism in response to mitochondrial metabolic dysfunction (38). Both, OctD and LCKD, significantly reduced the gene expression of the myomitoine fibroblast growth factor 21 in Cpt2<sup>−/−</sup> hearts, and LCKD also reduced growth differentiation factor 15 and glutathione-S-transferase alpha-1 (Fig. 3E, F). Thus, both ketogenic diets attenuated mitochondrial metabolic stress-induced myomitoine expression profiles in Cpt2<sup>−/−</sup> hearts, whereas minimal changes were observed in control hearts. Because medium-chain ketogenic diet corrected hypertrophy in a model of impaired pyruvate metabolism (21, 34), we next questioned the effect of the LCKD and OctD diets on regulators of pyruvate oxidation. Cpt2<sup>−/−</sup> hearts had increased pyruvate dehydrogenase kinase 4 mRNA abundance and PDH phosphorylation, and neither LCKD nor OctD altered these regulators of pyruvate oxidative metabolism in either genotype suggesting that the diets may not directly impact pyruvate oxidation (Fig. 3G–J). Together, these data suggest that LCKD, but not OctD, alleviates myomitoine expression but promotes further exacerbation of mTOR-growth pathway and pathological hypertrophy genes in CPT2-deficient hearts. Minimal effects on gene expression were observed in control mice, suggesting that short-term ketogenic diets rich in either long-chain or medium-chain fatty acids do not promote pathological hypertrophy or metabolic stress-induced myomitoine secretion in the normal heart.

**Cardiac acylcarnitine accumulation is not regulated by octanoate diet**

Cpt2<sup>−/−</sup> hearts accumulate significant amounts of long-chain acylcarnitines (LCACs) (23) on standard diet, so we questioned if the high octanoate diet would alleviate the acylcarnitine accumulation by increasing oxidative flux of carnitine-independent substrates (i.e., ketone bodies and octanoate). Surprisingly, OctD failed to correct the free carnitine deficiency or to attenuate LCAC accumulation in Cpt2<sup>−/−</sup> hearts (Fig. 4A, B). These data suggest that, despite diet-induced surplus of alternative substrates for oxidative metabolism, the steady-state levels of accumulated acylcarnitines in Cpt2<sup>−/−</sup> are not altered by OctD. LCAC accumulation and free carnitine deficiency are suspected regulators of cardiomyocyte function (39–42). However, in Cpt2<sup>−/−</sup> mice at end-stage disease and near heart failure, total LCACs did not increase, but several species decreased, including palmitoylcarnitine (Fig. 4C). Taken together, these data suggest that Cpt2<sup>−/−</sup> mice on standard low-fat diets reach the maximum capacity of LCAC accumulation that is not reduced by octanoate diet or altered by eminent heart failure. Because the provision of octanoate as alternative fuel did not restore free carnitine levels in Cpt2<sup>−/−</sup> mice, we thought to evaluate the link between carnitine deficiency and cardiac

ketogenic diet formulated with long-chain fatty acids as the primary fat source (long-chain fatty acid ketogenic diet [LCKD]) did neither alleviate hypertrophy nor prevented progression to heart failure in Cpt2<sup>−/−</sup> mice (23) when administered at 6 weeks of age and for a total of 4 weeks. Similarly, others have suggested that the failing heart increases ketone body utilization, and this could act as a potential contributor to failure (37). Here, we found that LCKD-fed Cpt2<sup>−/−</sup> hearts had increased expression of pathological remodeling markers, namely fibroblast growth factor 6, myosin heavy chain beta, and atrial natriuretic factor (Fig. 3C). LCKD significantly increased mTor gene expression but did not significantly increase p70S6K phosphorylation in Cpt2<sup>−/−</sup> hearts (Fig. 3D and supplemental Fig. S1B). These data suggest that LCKD, but not OctD, further exacerbates pathological remodeling in Cpt2<sup>−/−</sup> hearts.

![Fig. 2.](https://example.com/fig2.png)

Dietary octanoate did not improve heart function in Cpt2<sup>−/−</sup> mice and increased cardiac output in control mice. (A) Left ventricular mass relative to body weight, (B) flow velocity, and (C–E) cardiac functional parameters derived from 2D echocardiogram and Doppler analysis of control and Cpt2<sup>−/−</sup> mice on control or OctD, females, n = 3–4. Data are presented as mean ± SEM. Statistical analysis by two-way ANOVA. Means depicting a different letter indicate significant differences between groups (P ≤ 0.05). Cpt2, carnitine palmitoyltransferase 2; EDV, end-diastolic volume; PSV, peak-systolic volume; SV, stroke volume.
Fig. 3. Effects of long- and medium-chain ketogenic diets on cardiac remodeling and metabolic programs. A–F: Cardiac mRNA abundance normalized to control group on control diet (CD) of pathological hypertrophy markers, mTOR pathway, and myokines from control and Cpt2<sup>M−/−</sup> mice fed either CD, OctD, or LCKD, females, n = 4–6. G, H: Phosphorylation levels of PDH enzyme at Serine293 and mRNA abundance of PDH kinase (Pdk4) as regulators of pyruvate metabolism in hearts from control and Cpt2<sup>M−/−</sup> mice fed either CD, OctD, or LCKD, females, n = 4–6. Statistical analysis by 2-way ANOVA. Means depicting a different letter indicate significant differences between groups (P ≤ 0.05). Cpt2, carnitine palmitoyltransferase 2; LCKD, long-chain fatty acid ketogenic diet; mTOR, mechanistic target of rapamycin; OctD, octanoate diet; PDH, pyruvate dehydrogenase; Pdk4, pyruvate dehydrogenase kinase 4.
hypertrophy by supplementing mice with free carnitine in the drinking water for 4 weeks, as previously described and validated (25). Free carnitine supplementation failed to rescue cardiac hypertrophy or to modulate gene expression of pyruvate dehydrogenase kinase 4, pathological remodeling markers, mTor, and myomitis in Cpt2M−/− mice fed control or OctD, n = 3. Data are presented as mean ± SEM, all from female mice. Statistical analysis by 2-way ANOVA. Means depicting a different letter indicate significant differences between groups (P ≤ 0.05). *by genotype, #by diet among controls, and $by diet among Cpt2M−/−. Cpt2, carnitine palmitoyltransferase 2; DBs, double bonds; wo, weeks of.

CPT2-deficient hearts have reduced highly unsaturated fatty acids in phospholipids and are resistant to octanoate diet-induced membrane desaturation

To determine if LCACs could flux back through the reversible reaction catalyzed by CPTI into long-chain acyl-CoAs and be incorporated into complex lipids, the acyl chain length and saturation of cardiac membrane lipids were analyzed. Saturated acyl chains, the major substrates for mitochondrial oxidation, were reduced in Cpt2M−/− heart phospholipids suggesting
that acyl chains not used for oxidation are not fluxing into phospholipids. In addition, most of the phosphatidylcholine (PC) species containing ≥4 unsaturated bonds were depleted by 30–90%, whereas those containing 2 or 3 unsaturated bonds were increased by ~5- and 2-fold, respectively, in Cpt2M−/− hearts (Figs. 4E, F and supplemental Fig. S2). The OctD elicited changes in control hearts by reducing the membrane content of PCs with -6 or more unsaturated bonds by ~50% and by increasing PCs of -4 and -5 unsaturated bonds by ~80% (Figs. 4E, F and supplemental Fig. S2). The Cpt2M−/− mice, unlike controls, were resistant to diet-induced alterations in phospholipid acyl chain composition (Figs. 4E, F and supplemental Fig. S2). Together, these data demonstrate that loss of CPT2 resulted in an overall reduction in highly unsaturated fatty acids within cardiac membrane phospholipids, an outcome that was not altered by the high octanoate diet, whereas control mice reduced membrane unsaturation in response to the high octanoate diet.

Free octanoate is an oxidative substrate in liver but not in heart or skeletal muscle

Given the inability of the OctD to rescue Cpt2M−/− cardiomyopathy, we reasoned that this could relate to intrinsic deficiencies in octanoate uptake, activation, and/or oxidation in cardiac mitochondria. Therefore, oxidative capacity of octanoate between organs was determined in whole-tissue homogenates and isolated mitochondria from liver, heart, and skeletal muscle. For these experiments, the substrate of interest was added along with malate, to support flux through the tricarboxylic acid cycle, and with ADP, to stimulate respiration. In line with the notion that medium-chain fatty acids are oxidized independent of the carnitine shuttle, liver homogenates and isolated mitochondria were fully capable of oxidizing free octanoate in the absence of carnitine (Fig. 5A and supplemental Fig. S3). On the contrary, cardiac and skeletal muscle homogenates and isolated mitochondria showed no appreciable oxidation rates in the presence of free octanoate (Fig. 5A and supplemental Fig. S3). None of the tissues could oxidize the long-chain fatty acid palmitate in the absence of carnitine (Fig. 5B and supplemental Fig. S3) showing that the differential oxidative capacity between liver, heart, and skeletal muscle is chain length dependent. We next questioned why the liver, but not the heart or skeletal muscle, could oxidize free octanoate. Octanoate oxidation through the beta-oxidation spiral is initiated by the matrix enzyme MCAD whose substrate is octanoyl-CoA within the mitochondrial matrix. Octanoyl-CoA can be the product of either CPT2, utilizing CoA and octanoylcarnitine as substrate, or mitochondrial ACSMs, utilizing CoA and free octanoate as substrate. The liver, but not the heart or skeletal muscle, highly expressed several Acsm genes, namely Acsm1, Acsm3, and Acsm5 (Fig. 5C), as we have previously reported (43). Expression of ACSM3 protein was confirmed in liver mitochondria but could not be detected in skeletal muscle or heart (Fig. 5D and supplemental Fig. S3). Thus, the liver may have the unique ability to activate free octanoate for betaoxidation via ACSM. These data demonstrate that liver mitochondria, but not muscle or heart, can oxidize free octanoate.

Octanoylcarnitine is oxidized by liver, heart, and muscle in a CPT2-independent manner

We next assessed the ability of the tissues to use the acylcarnitine shuttle for oxidation of octanoate. Therefore, to represent the sequential substrates needed for the acylcarnitine shuttle reactions, free octanoate was used as the oxidative substrate, then CoA was added followed by free carnitine addition. Liver-isolated mitochondria and homogenates showed high rates of oxidation for free octanoate that were not further increased by addition of CoA or carnitine (Fig. 6A, C and supplemental Fig. S3). Conversely, heart mitochondria and homogenates only oxidized free octanoate after both, CoA and carnitine, were added to the reaction buffer (Fig. 6A, C and supplemental Fig. S3). To confirm that the lack of exogenous ATP in our assay conditions was not limiting the matrix ATP supply for ACSM-mediated reactions, an ATP-supplemented buffer was used and again showed high rates of free octanoate oxidation in liver, but not in heart (supplemental Fig. S3D–G). Control experiments were performed using palmitate as substrate to show that liver and heart failed to oxidize free palmitate until both CoA and carnitine were added in agreement with the dependence of long-chain fatty acids on the acylcarnitine shuttle for oxidation (Fig. 6B, C and supplemental Fig. S3). To assess if the lack of ACSMs in the heart limited octanoate oxidation, octanoyl-CoA, the product of ACSM action and substrate for MCAD, was provided as substrate. In liver, rates of octanoyl-CoA oxidation were high and nearly identical to those of free octanoate; in heart, however, octanoyl-CoA was not oxidized (Fig. 6D, E). To determine if octanoyl-CoA was not used by heart because of inability to be transported into the mitochondria, carnitine was added after octanoyl-CoA to allow production of octanoylcarnitine by CPT1 for transport into the mitochondria. Carnitine addition after octanoyl-CoA initiated a slow but steady increase in oxidation rates in the heart (Fig. 6D, E) suggesting that octanoyl-CoA is a viable substrate for the heart but only in the presence of free carnitine. We next questioned if the metabolic signature of skeletal muscle was similar to that of heart, and indeed skeletal muscle oxidation rates closely mimicked those observed in heart (Fig. 6D, E). Control experiments using palmitoyl-CoA followed by free carnitine addition confirmed that liver, heart, and skeletal muscle oxidize long-chain fatty acids only in the presence of carnitine (Fig. 6F). As for maximum
respiration capacity, heart and skeletal muscle mitochondria showed appreciably higher oxidative rates for both octanoate and palmitate when CoA and carnitine were subsequently added, compared with liver (Fig. 6E, F). To test respiratory rates when carnitine esters are provided as fuel, octanoylcarnitine was administered to show ∼4-fold greater oxidative rates in muscles compared with liver (Fig. 6G, H). In agreement, the heart presents higher protein abundance and has higher enzymatic activity for MCAD, a mitochondrial matrix dehydrogenase that could potentially set the rate for medium-chain fatty acid beta-oxidation (Fig. 6I). When considering long-chain fatty acid oxidation of palmitoylcarnitine, the heart and muscle had ∼9- and 7-fold greater rates than liver, respectively (Fig. 6J), suggesting that muscle tissues have overall high mFAO capacity.

Because heart and skeletal muscle had higher oxidation rates for octanoyl-CoA + Carn and octanoylcarnitine, and because the only known mitochondrial acyl-CoA-acylcarnitine exchanger is the carnitine shuttle (CPT1-CACT-CPT2), we decided to determine if carnitine-palmitoyltransferase activity is required for oxidation of octanoylcarnitine. Oxidation rates were determined in skeletal muscle mitochondria from control and skeletal muscle-specific CPT2-deficient mice (Cpt2Sk−/−) (24). In agreement with our previous findings in Cpt2M−/− hearts (23), the oxidation rates of octanoylcarnitine in skeletal muscle mitochondria were not different between control and Cpt2Sk−/− (Fig. 6K, L). Confirmation of CPT2 deficiency was evident by the nearly absent rates of palmitoylcarnitine oxidation in Cpt2Sk−/− muscle (Fig. 6K, L). Next, we assessed the involvement of
Fig. 6. Octanoylcarnitine is oxidized by liver, heart, and muscle in a CPT2-independent manner. A, B: Rates of oxygen consumption in isolated mitochondria of liver and heart as representative trace over time and (C) quantitation of maximum rates during administration of malate (M) and ADP with octanoate (Oct) or palmitate (PA) as substrates, followed by CoA and free L-carnitine (Carn), n = 3. D: Rates of oxygen consumption in isolated mitochondria from liver, heart, and skeletal muscle as representative trace and (E, F) quantitation of maximum rates when given M and ADP with octanoyl-CoA (OCoA) or palmitoyl-CoA followed by carnitine, n = 3. G: Rates of oxygen consumption in isolated mitochondria of liver, heart, and skeletal muscle as representative trace and (H) quantitation of maximum rates given octanoylcarnitine (OCarn) as substrate, n = 3. I: MCAD protein and activity in heart and liver, n = 3. J: Rates of palmitoylcarnitine oxidation in isolated mitochondria of liver, heart, and skeletal muscle. K: Representative trace of oxygen consumption and (L) quantification of maximum rates in isolated mitochondria of skeletal muscle from control or Cpt2Sk−/− mice given octanoylcarnitine and palmitoylcarnitine (PCarn), n = 3. M: Relative carnitine O-octanoyltransferase (CrOT) abundance in isolated mitochondria from liver and heart as detected by discovery proteomics, n = 5. Data are presented as mean ± SEM, all from male mice. *P ≤ 0.05 by one-way ANOVA (E, F, H, J, and M) or Student’s t-test (I and J). Means depicting a different letter indicate significant differences between groups (P ≤ 0.05). MCAD, medium-chain acyl-CoA dehydrogenase.
CrOT, a peroxisomal enzyme that performs the same reaction as CPT2 but has very high preference for octanoyl-CoA as substrate. We detected CrOT in mitochondrial fractions from both liver and heart, with higher abundance in liver when normalized to total proteins discovered (Fig. 6M, left panel). When correcting CrOT enrichment relative to peroxisomal proteins, CrOT was found at similar levels in the liver and heart (Fig. 6M, right panel). While it is not clear if CrOT is present in the mitochondria, these data suggest the possibility of CrOT activity replacing CPT2 for matrix octanoyl-CoA formation and subsequent oxidative metabolism. All together, these data suggest that the liver, heart, and muscle comprise the necessary machinery to oxidize octanoate in a carnitine-dependent, but CPT2-independent, manner, and that liver is capable of CoA- and carnitine-independent octanoate oxidation (Fig. 7).

DISCUSSION

We have previously reported that mediators of cardiac hypertrophy in Cpt2−/− mice are distinct from that of other common models of hypertrophy (23). Specifically, we have shown that unlike hypertension-induced and ischemia-induced models (36, 44–47), the cardiac hypertrophy triggered by CPT2 deficiency is resistant to attenuation by rapamycin, an mTor inhibitor, and by trichostatin A, a deacetylase inhibitor (23). Because both rapamycin and trichostatin A improve mitochondrial oxidative metabolism in normal hearts (48–53), our data suggested that restoration of cardiac fatty acid oxidation is required for the effectiveness of these therapies. In agreement, both high-fat diet and medium-chain ketogenic diet were shown to improve severe cardiac hypertrophy in models of impaired cardiac pyruvate oxidation (21, 34). These results suggest that increased flux of either ketones or fatty acids through oxidative metabolism can compensate for the loss of carbohydrate-oxidative metabolism and restore cardiac structure. Ketogenic diets have also been documented to paradoxically reduce ketolytic activity in the heart and increase long-chain fatty acid oxidative metabolism (54). Our data herein unequivocally show that in the absence of long-chain fatty acid oxidative flux in the mitochondria, medium-chain fatty acid diets or ketogenic diets cannot rescue cardiac hypertrophy. Thus, these combined data suggest that the ability of ketogenic diets to increase carnitine-dependent fatty acid oxidation in the heart is facilitating cardioprotective effects and that without long-chain fatty acid oxidative metabolism, cardioprotection cannot occur. This is surprising considering that medium-chain oil therapy is a standard of care for patients with long-chain fatty acid oxidation disorders (55, 56).

The ability of medium-chain fatty acids of eight carbons or less to produce ketone bodies following an oral bolus suggests that these fatty acids are preferred substrates for liver metabolism. Indeed, medium-chain fatty acids are well known to be digestively absorbed into the portal vein for direct transit to the liver. Thus, dietary medium-chain fatty acids bypass absorptive processing of common dietary fats (i.e., long-chain fatty acids), which are packaged into lipoprotein particles by enterocytes and released into the lymphatics, which merge into the vascular system near the heart. Thereby, while dietary long-chain fatty acids are first presented to the heart and then travel throughout the body passing through skeletal muscle before gaining access to the liver, dietary medium-chain fatty acids are directly routed from the gut to the liver. The liver is rich in the enzymatic machinery required for medium-chain fatty acid oxidation and is even capable of oxidizing free octanoate. Hepatocytes also incorporate octanoate into triglycerides for secretion into the circulation within lipoprotein particles (57). Therefore, heart and muscle may access majority of dietary...
octanoyl-CoA that medium-chain fatty acids are oxidized independent of carnitine. Thus, mammals may be biologically adapted to metabolize free octanoate and octanoylcarnitine in liver but only octanoylcarnitine in muscles.

To further delineate the mechanisms by which liver oxidizes free octanoate but cardiac and skeletal muscles cannot, we considered the fact that octanoate must be converted into octanoyl-CoA within the mitochondrial matrix for subsequent oxidative metabolism. Acyl-CoA moieties can be formed in the matrix by the transferase action of CPT2 when acylcarnitine and CoA are substrates or by the ligation action of ACSMs when free fatty acids and CoA are provided. Here, we demonstrate a striking distinction between liver and muscles for mitochondrial expression of ACSMs. The liver has high abundance of three of the five known ACSMs while ACSM expression is nearly absent in the heart and skeletal muscle (43). The action of these ASCMs may equip the liver with a unique advantage to activate and oxidize octanoate in the form of octanoyl carnitine, as a product of peroxisomal oxidation and peroxisomal CrOT enzyme action. Thus, mammals may be biologically adapted to metabolize free octanoate and octanoylcarnitine in liver but only octanoylcarnitine in muscles.

Acylcarnitines, short to long, cannot freely cross membranes, unlike their free fatty acid counterparts (58). Therefore, acylcarnitines are transported between cellular compartments, and in and out of cells, through protein-mediated processes. CACT is the best characterized example of a transporter of LCACs, located in the inner mitochondrial membrane. Indeed, patients with defects in CACT have identical clinical presentation to those with CPT2 deficiency (59). While transport of free octanoate into the matrix does not require a transporter, its esters, octanoylcarnitine and octanoyl-CoA, theoretically should require transporters because of their more hydrophilic nature. Intriguingly, CACT-deficient leukocytes cannot oxidize palmitate but are able to oxidize octanoate (60). We also found that CPT2-deficient hearts and muscles are fully capable of oxidizing octanoylcarnitine similar to our results herein using mouse CPT2-deficient muscles. The mitochondrial transporter used to facilitate octanoylcarnitine oxidation remains unestablished. Curiously, we found that liver presented with an additional, potentially transport-mediated, advantage for oxidation of octanoyl-CoA, which was readily oxidized by liver, but not muscles, in a manner independent of carnitine. These data suggest that either liver expresses mitochondrial transporters for octanoyl-CoA itself or that the CoA is cleaved from octanoate prior to entry into the mitochondrial matrix by acyl-CoA thioesterases that are active on medium-chain acyl-CoAs and abundant in liver (43, 61). Because our data for liver octanoyl-CoA oxidation are identical between homogenates and isolated mitochondria, we would argue that matrix thioesterases (62) are not involved, unless the formation of octanoyl-CoA within the matrix is not required for mitochondrial octanoyl-CoA oxidative metabolism. Given our data showing the rapid rise to identical rates of oxidation in liver for all three forms of octanoate, the free, -CoA, and -carnitine esters, we suggest that the transport-mediated processes and/or enzymatic metabolism of octanoate and its metabolites occur in a rapid and unrestrictive fashion in liver. Previous work demonstrated that inhibition of CPT1 with etomoxir or malonyl-CoA in muscle mitochondria does not affect octanoylcarnitine oxidation, suggesting that octanoylcarnitine oxidation occurs independent of CPT1 in muscle (63). Importantly, our findings using a CPT2 deficiency model show that CPT2 is not involved in the oxidation of octanoylcarnitine in muscles, yet the transporter and transferase responsible for octanoylcarnitine transport into the mitochondrial matrix and for its oxidative metabolism remain undefined. A possibility is the presence of CrOT or a CrOT-like enzyme in the mitochondrial matrix where it could convert octanoylcarnitine to octanoyl-CoA. Indeed, we identified CrOT protein in mitochondrial preparations of both liver and heart, thus there is a potential for CrOT to contribute to octanoylcarnitine oxidation in both tissues; however, its localization to the matrix has not been established.

It is not clear the exact chemical form and location that acylcarnitines take when accumulating inside cells. The amphipathic nature of acylcarnitines could hypothetically allow accumulation as micellar structures and/or be integrated into the amphipathic layers within membranes. Membrane accumulation of LCACs could increase packing defects, causing increased space between phospholipids. Packing defects are also caused by excessive changes to the ratio of saturated:unsaturated fatty acids within membranes (64, 65). This biophysical property is important for membrane curvature and protein-membrane interactions. It is possible that the reduction in highly unsaturated phospholipid species in Cpt2<sup>2/−</sup> hearts, and in Cpt2<sup>2/−</sup> muscles (24), is a compensatory mechanism to maintain membrane dynamics because of membrane acylcarnitine integration. Reduced membrane unsaturation is also reported in individuals with genetic defects in mFAO, and these patients are often supplemented with dietary DHA (66).
The OctD affected cardiac function of control mice, resulting in increased cardiac output. The increase in cardiac output could be the result of increased peripheral blood perfusion demands in response to increased oxygen requirement for high rates of oxidative metabolism during ketotic dietary conditions. However, the Cpt2M−/− mice were impervious to these diet-induced effects and maintained similar cardiac structural dimensions and functional outputs between the two diets indicating that metabolic remodeling triggered by CPT2 deficiency occurs prior to dietary manipulation and was not further changed by octanoate provision. Increasing the fatty acid chain length by one increment to decanoic acid necessitates the requirement of the carnitine shuttle for oxidation (67); therefore, patients with defects in long-chain mFAO should be supplemented with carefully curated medium-chain oils containing eight carbons or lower (68). Our data also suggest that while the liver does not require carnitine for octanoate oxidation, muscles do require carnitine. Because patients with fatty acid oxidation disorders present with free carnitine deficiency, the use of free carnitine supplementation is sometimes implemented; however, carnitine supplementation remains debated because of concerns that it facilitates cardiac LCAC accumulation and subsequent cardiac arrhythmias (66). Our data herein suggest that free carnitine supplementation along with medium-chain therapy could facilitate the oxidation of medium-chain fatty acid oxidation in heart and muscle tissues.

Together, our findings demonstrate that liver mitochondria have the ability to oxidize free octanoate and octanoyl-CoA in the absence of carnitine, whereas muscle and heart mitochondria do not. However, liver, heart, and muscles oxidize octanoylcarnitine, independent of CPT2. We speculate that this differential oxidative capacity is a biological adaptation to the manner in which octanoate and its metabolites are presented to the mitochondria within these tissues. Specifically, the liver could be predominantly exposed to the free form of octanoate from a dietary source during intestinal absorption, as well as, from peroxisomal oxidation, whereas muscles are more exclusively exposed to octanoylcarnitine as a product of peroxisomal oxidation. Our work also suggests that, as part of such biological adaptations, the liver and heart express distinct enzymatic machinery for octanoate oxidation, such as ACSMs (Fig. 7). Finally, failure of octanoate ketogenic diet to restore normal cardiac structure and function suggests that oxidation of long-chain fatty acids is critical and irreplaceable for cardiac health.

Data availability
All the data are contained within the article.  

Supplemental data
This article contains supplemental data.
and obesity: only a matter of fat? High Blood Press. Cardiometab. Res. 22, 29–41
6. Cho, I. J., Chang, H. J., Park, H. B., Heo, R., Shin, S., Shim, G. Y., Hong, G. R., and Chung, N. (2015) Aortic calcification is associated with arterial stiffening, left ventricular hypertrophy, and diastolic dysfunction in elderly male patients with hypertension. J. Hypertens. 33, 1033–1041
7. Leveil, E., Mahmood, M., Piechnik, S. K., Ariga, R., Francis, J. M., Rodgers, C. T., Clarke, W. T., Sabharwal, N., Schneider, J. E., Karamitsos, T. D., Clarke, K., Rider, O. J., and Neubauer, S. (2016) Relationship Between Left Ventricular Structural and Metabolic Remodeling in Type 2 Diabetes. Diabetes. 65, 44–52
8. Levy, D., Anderson, K., Savage, D., Kannel, W., Christiansen, J., and Castelli, W. (1988) Echocardiographically detected left ventricular hypertrophy: prevalence and risk factors. The Framingham Heart Study. Am. Intern. Med. 108, 7–13
9. Ingwall, J. S. (2009) Energy metabolism in heart failure and remodelling. Cardiovasc. Res. 81, 412–419
10. Ventura-Clapier, R., Garrier, A., and Viskers, V. (2004) Energy metabolism in heart failure. J. Physiol. 555, 1–13
11. Lehman, J. J., and Kelly, D. P. (2002) Gene regulatory mechanisms governing energy metabolism during cardiac hypertrophic growth. Heart Fail. Rev. 7, 173–185
12. Gibb, A. A., Gilbert, L. A., Lorkiewicz, P., Trainor, P., Tseng, M. T., Enright, P. N., Bhatnagar, A., and Hill, B. G. (2016) Abstract 1044A: altered cardiomyocyte glycosylation causes cardiac hypertrophy and mitochondrial dysfunction. Circulation. 134 (Suppl 1), A19448-A
13. Ellis, J. M., Mentock, S. M., DePetriolo, M. A., Koves, T. R., Sen, S., Watkins, S. M., Muoio, D. M., Cline, G. W., Taegtmeyer, H., Shima, D. T., Willis, M. S., and Coleman, R. A. (2011) Mouse cardiac acyl coenzyme A synthetase 1 deficiency impairs fatty acid oxidation and induces cardiac hypertrophy. Mol. Cell Biol. 31, 1252–1262
14. Haynie, K. R., Vandanmagsar, B., Wicks, S. E., Zhang, J., and Haynie, K. R. (2015) Mitochondrion as a target for heart failure: a comprehensive review. Front. Physiol. 6, 46
15. McCombs, K. S., Kovacs, A., Weinheimer, C. J., Shew, T. M., Koves, T. R., Ilkayeva, O., Kamph, D. R., Pyles, K. D., King, M. T., Veech, R. L., DeBosch, B. J., Muoio, D. M., Gross, R. W., and Finck, B. N. (2020) Ketogenic diet prevents heart failure from defective mitochondrial pyruvate metabolism. bioRxiv. https://doi.org/10.1101/2020.02.21.959635
16. Salubrancamian, S., Johnston, R. K., Moschella, P. C., Mani, S. K., Tuxworth, W. J., Jr., and Kuppuswamy, D. (2009) mTOR in growth and protection of hypertrophying myocardium. Cardiovasc. Hematol. Agents Med. Chem. 7, 52–63
17. McMullen, J. R., Sherwood, M. C., Tarnavski, O., Zhang, L., Dorfman, A. L., Shiio, T., and Izumo, S. (2004) Inhibition of mTOR signaling with rapamycin regresses established cardiac hypertrophy induced by pressure overload. Circulation. 109, 3050–3055
18. Aubert, G., Martin, O. J., Horton, J. L., Lai, L., Vega, R. B., Leone, T. C., Koves, T., Gardell, S. J., Kristin, M., Hoppel, C. L., Lewandowski, E. D., Crawford, P. A., Muoio, D. M., and Kelly, D. P. (2016) The failing heart relies on ketone bodies as a fuel. Circulation. 133, 679–705
19. Kliewer, S. A., and Mangelsdorf, D. J. (2019) A dozen years of discovery: insights into the physiology and pharmacology of FGF21. Cell Metab. 20, 246–253
20. Koca, J., Makreka-Kuka, M., Cirule, H., Grinberga, S., Sevostjana, E., Dambrova, M., and Liepinsh, E. (2017) Decrease in long-chain acylcarnitine tissue content determines the duration of and correlates with the cardioprotective effect of methyl-GBB. Basic Clin. Pharmacol. Toxicol. 121, 106–112
21. Llobet, G., Gópala, K., and Jee, J. E. (2016) Lipotoxicity in obesity and diabetes-related cardiac dysfunction. Biochem. Biophys. Acta Mol. Cell Biol. Lipids. 1861, 1555–1568
41. Liepins, E., Makreka-Kuka, M., Volska, K., Kuka, J., Makarova, E., Antone, U., Sevostjanov, E., Vilskerst, R., Strods, A., Taras, K., and Dambrova, M. (2016) Long-chain acylcarnitines determine ischaemia/reperfusion-induced damage in heart mitochondria. Biochem. J. 473, 1193–1202

42. Yamada, K. A., Kanter, E. M., and Newatia, A. (2000) Long-chain acylcarnitine induces Ca2+ efflux from the sarcoplasmic reticulum. J. Cardiovasc. Pharmacol. 36, 14–21

43. Ellis, J. M., Bowman, C. E., and Wolfgang, M. J. (2015) Metabolic and tissue-specific regulation of acyl-CoA metabolism. PLoS One. 10, e0116587

44. Morales, C. R., Li, D. L., Pedrozo, Z., May, H. I., Jiang, N., Kyrck, S., Abel, E. D., Symons, J. D., and Jalili, T. (2009) Mammalian target of rapamycin is a critical regulator of cardiac hypertrophy in spontaneously hypertensive rats. Hypertension. 54, 1219–1227

45. Kuzman, J. A., O’Connell, T. D., and Gerdes, A. M. (2007) Rapamycin prevents thyroid hormone-induced cardiac hypertrophy. Endocrinology. 148, 3477–3484

46. Ooi, J. Y., Tuano, N. K., Rafehi, H., Ziemann, M., Du, X. J., and El-Osta, A. (2015) HDAC inhibition attenuates cardiac hypertrophy by acetylation and deacetylation of target genes. Epigenetics. 10, 418–430

47. Liepinsh, E., Makrecka-Kuka, M., Volska, K., Kuka, J., Makarova, E., Antone, U., Sevostjanov, E., Vilskerst, R., Strods, A., Taras, K., and Dambrova, M. (2016) Long-chain acylcarnitines determine ischaemia/reperfusion-induced damage in heart mitochondria. Biochem. J. 473, 1193–1202

48. Bonnefont, J. P., Demaille, G., Coudert, C., Demuynck, J., Brivet, M., Abadi, N., and Thuillier, L. (1999) Carnitine palmitoyltransferase deficiencies. Mol. Genet. Metab. 68, 424–440

49. Wunderling, K., Leopold, C., Jamitzy, I., Yaghmouir, M., Zink, F., Kratky, D., and Thiele, C. (2020) Hepatic synthesis of triacylglycerols containing medium-chain fatty acids is dominated by diacylglycerol acyltransferase 1 and efficiently inhibited by etomoxir. Mol. Metab. 10, 1–20

50. Haeyaert, P., Verdonck, A., and Van Gaal, F. H. (1987) Influence of acylcarnitines of different chain length on pure and mixed phospholipid vesicles and on sarcoplasmic reticulum vesicles. Chem. Phys. Lipids. 45, 49–63

51. Rubio-Gozalo, M. E., Bakker, J. A., Waterham, H. R., and Wers, R. J. (2004) Carnitine-acylcarnitine translocase deficiency, clinical, biochemical and genetic aspects. Mol. Aspects Med. 25, 521–532

52. Wunderling, K., Leopold, C., Jamitzy, I., Yaghmouir, M., Zink, F., Kratky, D., and Thiele, C. (1995) A patient with lethal cardiomyopathy and a carnitine-acylcarnitine translocase deficiency. J. Inherit. Metab. Dis. 18, 239–252

53. Tillander, V., Alexson, S. E. H., and Cohen, D. E. (2017) Deactivating fatty acids: Acyl-CoA thioesterase-mediated control of lipid metabolism. Trends Endocrinol. Metab. 28, 473–484

54. Bekeova, G., Anderson-Pullinger, L., Boye, K., Boos, F., Sharpsky, Y., Herrmann, J. M., and Seifert, E. L. (2019) Multiple mitochondrial thioesterases have distinct tissue and substrate specificity and CoA regulation, suggesting unique functional roles. J. Biol. Chem. 294, 19034–19047

55. Noland, R. C., Woodlief, T. L., Whitfield, B. R., Manning, S. M., Evans, J. R., Dudek, R. W., Lust, R. M., and Corrigan, R. N. (2007) Peroxisomal-mitochondrial oxidation in a rodent model of obesity-associated insulin resistance. Am. J. Physiol. Endocrinol. Metab. 293, E980–991

56. Harayama, T., and Riezman, H. (2018) Understanding the diversity of membrane lipid composition. Nat. Rev. Mol. Cell Biol. 19, 281–296

57. Manni, M. M., Tiberti, M. L., Pagnotta, S., Barelli, H., Gautier, R., Jacob, J., Beldi, F., Koyal, I., and Merino, B. (2015) Carnitine-acylcarnitine translocase deficiency, clinical, biochemical and genetic aspects. Mol. Aspects Med. 25, 521–532

58. Vitoria, L., Mestri-Hernández, E., Peña-Quintana, L., Bueno, M., Quijada-Fraile, P., Dalmau, J., Molina-Marrero, S., Pérez, B., and Merino, B. (2015) Carnitine-acylcarnitine translocase deficiency: experience with four cases in Spain and review of the literature. JIMD Rep. 20, 11–20

59. Wunderling, K., Leopold, C., Jamitzy, I., Yaghmouir, M., Zink, F., Kratky, D., and Thiele, C. (2020) Hepatic synthesis of triacylglycerols containing medium-chain fatty acids is dominated by diacylglycerol acyltransferase 1 and efficiently inhibited by etomoxir. Mol. Metab. 10, 1–20

60. Wunderling, K., Leopold, C., Jamitzy, I., Yaghmouir, M., Zink, F., Kratky, D., and Thiele, C. (1995) A patient with lethal cardiomyopathy and a carnitine-acylcarnitine translocase deficiency. J. Inherit. Metab. Dis. 18, 239–252

61. Tillander, V., Alexson, S. E. H., and Cohen, D. E. (2017) Deactivating fatty acids: Acyl-CoA thioesterase-mediated control of lipid metabolism. Trends Endocrinol. Metab. 28, 473–484

62. Bekeova, G., Anderson-Pullinger, L., Boye, K., Boos, F., Sharpsky, Y., Herrmann, J. M., and Seifert, E. L. (2019) Multiple mitochondrial thioesterases have distinct tissue and substrate specificity and CoA regulation, suggesting unique functional roles. J. Biol. Chem. 294, 19034–19047

63. Noland, R. C., Woodlief, T. L., Whitfield, B. R., Manning, S. M., Evans, J. R., Dudek, R. W., Lust, R. M., and Corrigan, R. N. (2007) Peroxisomal-mitochondrial oxidation in a rodent model of obesity-associated insulin resistance. Am. J. Physiol. Endocrinol. Metab. 293, E980–991

64. Harayama, T., and Riezman, H. (2018) Understanding the diversity of membrane lipid composition. Nat. Rev. Mol. Cell Biol. 19, 281–296

65. Manni, M. M., Tiberti, M. L., Pagnotta, S., Barelli, H., Gautier, R., Jacob, J., Beldi, F., Koyal, I., and Merino, B. (2015) Carnitine-acylcarnitine translocase deficiency, clinical, biochemical and genetic aspects. Mol. Aspects Med. 25, 521–532

66. Vitoria, L., Mestri-Hernández, E., Peña-Quintana, L., Bueno, M., Quijada-Fraile, P., Dalmau, J., Molina-Marrero, S., Pérez, B., and Merino, B. (2015) Carnitine-acylcarnitine translocase deficiency: experience with four cases in Spain and review of the literature. JIMD Rep. 20, 11–20

67. Chalmers, R. A., Stanley, C. A., English, N., and Wigglesworth, J. S. (1997) Mitochondrial carnitine-acylcarnitine translocase deficiency, clinical, biochemical and genetic aspects. Mol. Aspects Med. 25, 521–532

68. Roe, C. R., and Mochel, F. (2006) Anaplerotic diet therapy in inherited metabolic disease: ketogenic nutriend environment. J. Biol. Chem. 285, 24447–24456

69. Longo, N., Atam di San Filippo, C., and Pasquali, M. (2006) Disorders of carnitine transport and the carnitine cycle. Am. J. Med. Genet. C Semin. Med. Genet. 142c, 77–85

70. Bonnefont, J. P., Demaille, G., Coudert, C., Demuynck, J., Brivet, M., Abadi, N., and Thuillier, L. (1999) Carnitine palmitoyltransferase deficiencies. Mol. Genet. Metab. 68, 424–440

71. Chalmers, R. A., Stanley, C. A., English, N., and Wigglesworth, J. S. (1997) Mitochondrial carnitine-acylcarnitine translocase deficiency, clinical, biochemical and genetic aspects. Mol. Aspects Med. 25, 521–532

72. Roe, C. R., and Mochel, F. (2006) Anaplerotic diet therapy in inherited metabolic disease: ketogenic nutrient environment. J. Biol. Chem. 285, 24447–24456

Differential metabolism of octanoate by liver and muscles