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Exogenous dietary fat can induce obesity and promote diabetes, but endogenous fat production is not thought to affect skeletal muscle insulin resistance, an antecedent of metabolic disease. Unexpectedly, the lipogenic enzyme fatty acid synthase (FAS) was increased in the skeletal muscle of mice with diet-induced obesity and insulin resistance. Skeletal muscle–specific inactivation of FAS protected mice from insulin resistance without altering adiposity, specific inflammatory mediators of insulin signaling, or skeletal muscle levels of diacylglycerol or ceramide. Increased insulin sensitivity despite high-fat feeding was driven by activation of AMPK without affecting AMP content or the AMP/ATP ratio in resting skeletal muscle. AMPK was induced by elevated cytosolic calcium caused by impaired sarco/endoplasmic reticulum calcium ATPase (SERCA) activity due to altered phospholipid composition of the sarcoplasmic reticulum (SR), but came at the expense of decreased muscle strength. Thus, inhibition of skeletal muscle FAS prevents obesity-associated diabetes in mice, but also causes muscle weakness, which suggests that mammals have retained the capacity for lipogenesis in muscle to preserve physical performance in the setting of disrupted metabolic homeostasis.
Muscle lipogenesis balances insulin sensitivity and strength through calcium signaling

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Exogenous dietary fat can induce obesity and promote diabetes, but endogenous fat production is not thought to affect skeletal muscle insulin resistance, an antecedent of metabolic disease. Unexpectedly, the lipogenic enzyme fatty acid synthase (FAS) was increased in the skeletal muscle of mice with diet-induced obesity and insulin resistance. Skeletal muscle–specific inactivation of FAS protected mice from insulin resistance without altering adiposity, specific inflammatory mediators of insulin signaling, or skeletal muscle levels of diacylglycerol or ceramide. Increased insulin sensitivity despite high-fat feeding was driven by activation of AMPK without altering AMP content or the AMP/ATP ratio in resting skeletal muscle. AMPK was induced by elevated cytosolic calcium caused by impaired sarco/endoplasmic reticulum calcium ATPase (SERCA) activity due to altered phospholipid composition of the sarcoplasmic reticulum (SR), but came at the expense of decreased muscle strength. Thus, inhibition of skeletal muscle FAS prevents obesity-associated diabetes in mice, but also causes muscle weakness, which suggests that mammals have retained the capacity for lipogenesis in muscle to preserve physical performance in the setting of disrupted metabolic homeostasis.

Introduction

By 2050, 1 in 3 Americans may have obesity-related type 2 diabetes, an often lethal condition (1) characterized by insulin resistance (2, 3). Skeletal muscle insulin resistance accounts for approximately 90% of impaired glucose disposal in human type 2 diabetes (4), and abnormal lipid metabolism is implicated in this process (5). Manipulating genes affecting different steps in lipid handling produces varying effects on glucose metabolism that have been interpreted to reflect increased insulin sensitivity with decreased intracellular fatty acid content (6–11). Despite this apparently inverse relationship between lipid content and insulin sensitivity, there is no consensus regarding how fatty acid flux in muscle affects insulin sensitivity (12, 13). Unexplored is the possibility that endogenous lipid synthesis in skeletal muscle affects diabetes risk.

Fatty acid synthase (FAS) (14) uses products of the citric acid cycle to synthesize the saturated fatty acid palmitate, contributing to de novo lipogenesis. Skeletal muscle de novo lipogenesis is not thought to be a major contributor to total muscle fatty acid flux (15). However, several lines of evidence suggest that de novo lipogenesis may serve a signaling function independent of providing lipid storage (16). As one example, FAS generates an endogenous ligand for the nuclear receptor PPARα in liver (17). PPARα is also found in muscle, where its overexpression has been implicated in the pathogenesis of insulin resistance (18). These observations raise the possibility that fatty acid flux generated by skeletal muscle FAS could promote insulin resistance through activation of PPARα.

To test the hypothesis that FAS affects skeletal muscle insulin sensitivity, we selectively inactivated FAS in this tissue. Our findings suggest that mice deficient in de novo lipogenesis were protected from skeletal muscle insulin resistance induced by high-fat diet (HFD) feeding. However, this protection did not appear to be mediated by PPARα. Instead, FAS deficiency in muscle altered the phospholipid composition and transport functions of the sarcoplasmic reticulum (SR; a relative of the endoplasmic reticulum adapted to sequester calcium), leading to elevated cytosolic calcium concentrations. Elevated intracellular calcium increased glucose transport through AMPK, but this metabolic benefit came at the cost of decreasing physical performance. Improved insulin sensitivity appeared to be physiologically connected to muscle fatigue through increased cytosolic calcium, which activated AMPK but likely prevents appropriate relaxation of muscle fibers, thus limiting optimal subsequent contraction.

Results

Skeletal muscle FAS is associated with insulin resistance. 4 weeks of feeding HFD to control mice predictably increased levels of glucose, insulin, and other metabolites affected by insulin resistance (see below). This intervention is known to decrease FAS expression at major sites of lipogenesis, such as liver and white adipose tissue (19, 20). Surprisingly, HFD feeding did not suppress FAS in muscle: mRNA, protein, and enzyme activity of FAS were increased in skeletal muscle after 4 weeks of HFD feeding (Figure 1A and Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI65726S1). FAS protein abundance was greater in slow-twitch muscles that are susceptible to diet-induced insulin resistance (21), such as soleus (Figure 1B), which suggests that muscle FAS may promote insulin resistance. To address this possibility, we crossed mice bearing loxP sites in Fasn (22) with transgenic mice expressing Cre recombinase driven by the human α–skeletal actin promoter (23). This strategy yielded FAS knockout in skeletal muscle (FASKOS) mice.

FASKOS mice are protected from diet-induced skeletal muscle insulin resistance. FASKOS mice had deficient FAS expression in skeletal muscle and were protected from diet-induced diabetes (Figure 2A). To delineate the mechanism of this protection, we measured glucose and insulin concentrations; both were decreased in FASKOS mice, suggesting that FAS deficiency affects insulin action (Figure 2B). Imaging of muscle sections stained with C2C12 brightly labeled the myofibrillar component in control mice, whereas FASKOS mice showed a loss of myofibrillar organization with a more granular appearance (Figure 3A). Similarly, protein expression of α-actinin in FASKOS mice was lower than controls (Figure 3B). PPARα protein expression was also decreased in FASKOS mice (Figure 3B). These data suggest that FAS promotes insulin resistance through increased PPARα expression, which alters calcium homeostasis in muscle.

Conflict of interest: The authors have declared that no conflict of interest exists.

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muscle, but not in liver, heart, or pancreas (Figure 1, C and D, and Supplemental Figure 1B). There was no genotype-specific effect on body weight or composition, as determined by MRI, with chow or HFD feeding (Figure 2A). Tissue weights of skeletal muscles, liver, and adipose depots were unaffected in FASKOS mice (Figure 2, B and C). Serum levels of glucose, free fatty acids, triglycerides, and cholesterol as well as of insulin and leptin were higher in HFD-versus chow-fed animals, but these effects were the same in control and FASKOS mice (Figure 2, D and E). The total skeletal muscle content of triglycerides, free fatty acids, malonyl-CoA, diacylglycerol (DAG), ceramide, and other sphingolipid species did not differ between HFD-fed control and FASKOS mice (Figure 3, A–E, and data not shown). These negative results are consistent with previous reports of low levels of lipogenic genes, including FAS, in skeletal muscle compared with expression of malonyl-CoA decarboxylase (15), which suggests that FAS does not have major effects on the total mass of lipids and lipid precursors in muscle.

Muscle FAS deletion did not alter metabolic rate, as measured by indirect calorimetry (Figure 4A). Respiratory quotient (RQ; reflecting choice of carbohydrate or lipid for energy production, with a lower value indicating more lipid use) was decreased in HFD-fed control mice that were awake and eating (during the dark phase of a daily cycle; Figure 4B), suggestive of less postprandial carbohydrate metabolism consistent with HFD-induced insulin resistance. However, FASKOS mice were partially protected from this HFD-induced attenuation of RQ in the dark phase (Figure 4B), indicative of increased use of carbohydrates as fuel. To determine whether this preference for carbohydrate was due to increased insulin sensitivity, FASKOS mice were further metabolically phenotyped. Glucose and insulin tolerance tests showed that with chow feeding, FASKOS mice were indistinguishable from control mice (Figure 4, C and D). However, FASKOS mice were partially protected from HFD-induced glucose intolerance and insulin resistance (Figure 4, E and F). In hyperinsulinemic euglycemic clamp studies, the insulin-stimulated glucose disposal rate was 4-fold greater in FASKOS than in control mice, whereas there was no difference in the ability of insulin to suppress endogenous glucose production (mostly reflecting hepatic effects) or release of fatty acids from fat (Figure 5, A–E). The findings that increased glucose disposal occurred in the absence of differences in adiposity, potential circulating contributors to insulin resistance, or metabolic rate (Figures 2 and 4) suggest that muscle FAS deficiency increased insulin sensitivity through effects intrinsic to muscle.

Muscle FAS deletion results in intrinsic AMPK activation. Insulin-stimulated glucose uptake in skeletal muscle is mediated by translocation of GLUT4 to the cell surface (24). In muscles isolated after the clamp, GLUT4 abundance was not different between control and FASKOS mice (Figure 5F). There was also no effect of FAS deletion on phosphorylation of Akt (pAktThr308 and pAktSer473) or of its substrate, glycogen synthase kinase–3 (GSK3), in insulin-stimulated muscles, which suggests that these mediators of insulin receptor signaling are not involved in the FASKOS phenotype. However, phosphorylation of Akt substrate of 160 kDa (AS160), an Akt substrate Rab GTPase activating protein known to regulate GLUT4 uptake in skeletal muscle, was increased in FASKOS mice (Figure 5G). This suggests that FAS deficiency increased insulin sensitivity through effects intrinsic to muscle.
translocation (25, 26), was increased in the muscles from FASKOS compared with control mice (Figure 5F). AS160 phosphorylation is permissive for glucose transport. In addition to Akt, AMPK can phosphorylate AS160 (27, 28). Phosphorylation of AMPK and its substrate, acetyl-CoA carboxylase (ACC), were found to be elevated in FASKOS versus control mice (Figure 5F). Because AMPK enhances muscle insulin sensitivity, these results suggest that the insulin sensitivity phenotype in HFD FASKOS mice may be due to muscle-intrinsic AMPK activation.

Soleus muscles were isolated from control and FASKOS mice fed chow or HFD, then incubated with 2-deoxyglucose. As with intact animals, there was no effect of FAS inactivation in the chow condition, but soleus and extensor digitorum longus (EDL) muscles isolated from FASKOS mice were partially protected from the HFD-induced reduction of insulin-stimulated glucose uptake (Figure 6A and Supplemental Figure 2). Western blotting of these isolated muscles confirmed the findings of others that HFD decreased AMPK activation in control muscles (29, 30) and showed that FASKOS muscles were protected from this suppression of AMPK activation (reflected as phosphorylation of AMPK itself as well as of ACC; Figure 6, B and C). Mirroring the findings in intact mice, Akt activation was unaffected, whereas AS160 phosphorylation was increased, in muscle of HFD-fed FASKOS mice (Figure 6, D and E). These observations suggest that FASKOS muscle is protected from HFD-induced suppression of AMPK activity and, since HFD feeding increased muscle FAS activity (Figure 1A), that skeletal muscle FAS inhibits AMPK activation.

Muscle-intrinsic AMPK activation is mediated by increased cytosolic calcium. To determine whether regulation of AMPK by FAS can be modeled in a cell line, C2C12 myocytes were treated with a lentivirus expressing an FAS shRNA. C2C12 cells are a transformed cell line with metabolic needs differing from those of intact muscle, but there is precedent for the use of these cells to gain insight into...
processes relevant to the role of muscle in physiology (31). FAS knockdown (FASKD) had no effect on activation of Akt or GSK3 compared with control cells treated with a scrambled virus, but increased phosphorylation of AMPK as well as of its targets, ACC and AS160 (Figure 7A), replicating the effects in FASKOS skeletal muscle. FAS generates an endogenous ligand for the nuclear receptor PPAR\(\alpha\) in liver (17), and PPAR\(\alpha\) deletion activates AMPK in skeletal muscle (18), which raises the possibility that FAS deletion affects AMPK through PPAR\(\alpha\). However, FAS deletion did not affect PPAR\(\alpha\) luciferase activity, and treatment with the PPAR\(\alpha\) agonist WY14643 did not reverse FASKD-induced AMPK activation (Figure 7, B and C). AMPK can be allosterically activated by AMP, but AMP content and the AMP/ATP ratio were unaffected by FAS deletion (Supplemental Figure 3, A and B). Total skeletal muscle malonyl-CoA levels were not different between control and FASKOS mice (Figure 3C). It is conceivable that changes in malonyl-CoA limited to discrete subcellular compartments might activate AMPK by inhibiting carnitine palmitoyl transferase–1 and fatty acid oxidation. However, incubation of C2C12 myocytes with the fatty acid oxidation inhibitor etomoxir did not alter AMPK signaling (data not shown), and fatty acid oxidation rates were not different from controls in muscles or C2C12 cells with FAS deletion (Supplemental Figure 3, C and D). Inflammation can play a role in insulin sensitivity, but there was no effect of FAS deletion on JNK or IKK\(\beta\) activity in skeletal muscle (Supplemental Figure 4). There was also no detectable interaction between FAS and AMPK in co-IP experiments (data not shown). Therefore, the effect of muscle FAS deletion on AMPK activation does not appear to be mediated by PPAR\(\alpha\), AMP levels, altered fatty acid oxidation, inflammation, or a direct effect of FAS on AMPK.

AMPK can be phosphorylated by liver kinase B1 (LKB1) or Ca\(^{2+}\)/calmodulin-dependent kinase kinase–\(\beta\) (CaMKK\(\beta\)). FASKD did not affect mRNA levels or protein abundance of LKB1, CaMKK\(\alpha\), or CaMKK\(\beta\) (Supplemental Figure 5). LKB1 appears to be constitutively active, but CaMKK\(\beta\) is activated by increased cytosolic calcium (32). Since FAS inactivation in cardiac muscle increases calcium signaling (33), and calcium (34), calmodulin (35), and CaMKII (36) have been implicated in glucose uptake, we addressed a potential role for calcium. In C2C12 myocytes, cytosolic calcium content increased in response to FASKD (Figure 7, D and E). The increase in cytosolic calcium triggered the Ca\(^{2+}\)/calmodulin signaling cascade in C2C12 myocytes (Figure 8, A and B). Activation of CaMKK\(\beta\) (detected as CaMKI phosphorylation) was also found in clamped as well as isolated FASKOS muscles (Figure 8, C and D). To determine whether CaMKK\(\beta\) was required for the effect of FAS deletion on AMPK activation, we inhibited CaMKK\(\beta\) in the setting of FASKD in C2C12 myocytes. FASKD-induced AMPK activation was abrogated by CaMKK\(\beta\) knockdown, which was confirmed by treatment with a chemical inhibitor of CaMKK, STO-609 (Figure 8, E and F). These data suggest that induction of AMPK activity with FAS deletion is related to increased calcium signaling.

Muscle FAS affects SERCA by altering SR phospholipid composition. Control of cytosolic calcium in skeletal muscle is mediated by transporters found at the plasma membrane and at the SR. Dogma holds that FAS is exclusively cytoplasmic, but subcellular fractionation studies of skeletal muscle unexpectedly showed that, in addition to the cytoplasm, SR was also highly enriched in FAS (Figure 9A). To address the possibility that FAS physically interacts with SR proteins to regulate calcium flux, the SR fraction from mouse hindlimb muscles was treated with an FAS antibody, and the immunoprecipitated proteins were identified by
SERCA2 was palmitoylated in sensitive assays involving replacement of biotin for palmitate at cysteine residues (data not shown). However, neither SERCA1 nor SERCA2 was palmitoylated in sensitive assays involving replacement of biotin for palmitate at cysteine residues (data not shown).

To determine whether the lipid environment of the SR is altered by FAS deficiency, the SR fraction of hindlimb muscle was isolated and analyzed by mass spectrometry (Supplemental Table 1). It is likely that several of these proteins do not reside in the SR, given the sensitivity of mass spectrometry and possible contamination of the SR fraction with proteins from other compartments. However, 3 isoforms of sarco/endoplasmic reticulum calcium ATPase (SERCA), the ATPase that transports cytosolic calcium to the SR lumen, were detected. Results of co-IP experiments using SERCA1 and SERCA2 antibodies were consistent with an association between these 2 proteins and FAS (Figure 9B). ACC did not appear to interact with the SERCAs (Figure 9B). Immunofluorescence colocalization studies in C2C12 cells also suggested an interaction between FAS and SERCA1 (Figure 9C). FAS also colocalized with the endoplasmic reticulum/SR marker calnexin, but not with the Golgi marker GM130 (Figure 9C). FASKD in C2C12 myocytes decreased SERCA enzyme activity without affecting SERCA1 protein abundance (Figure 9D). Transfecting C2C12 myocytes with a vector expressing myc-tagged SERCA1 restored SERCA enzyme activity (Figure 9D). Expressing myc-tagged SERCA1 also eliminated the FASKD-induced increase in AMPK phosphorylation (Figure 9E), consistent with the concept that FAS inactivation impairs SERCA activity to activate AMPK. FASKD did not affect myc-tagged SERCA1 expression (Figure 9E). In addition, SERCA enzyme activity was decreased in soleus and EDL muscle of HFD-fed, but not chow-fed, FASKOS mice without affecting SERCA1 protein abundance (Figure 9F and Figure 10A). Transfecting C2C12 myocytes with a vector expressing myc-tagged SERCA1 restored SERCA enzyme activity (Figure 9F). Transfecting C2C12 myocytes with a vector expressing myc-tagged SERCA1 restored SERCA enzyme activity (Figure 9F).
Discussion

Synthesizing fatty acids in skeletal muscle would not be predicted to be relevant to HFD-induced insulin resistance. Since FAS is decreased by exogenous fat in lipogenic tissues (19, 20), one might also anticipate that excess dietary fat would decrease FAS in muscle. Unexpectedly, we found that HFD feeding increased FAS activity in skeletal muscle of control mice in concert with the induction of insulin resistance. FASKOS mice, with skeletal muscle–specific inactivation of FAS, were partially protected from diet-induced insulin resistance, but also prone to diet-induced muscle weakness. The underlying mechanism involves an effect of FAS on the modulation of cytosolic calcium. We found that FAS, thought to be exclusively cytosolic, was also present in the SR fraction of skeletal muscle, where it is physically associated with SERCA proteins. FAS deficiency decreased SERCA activity by altering the phospholipid composition of the SR, which resulted in increased cytosolic calcium. Decreased uptake of cytosolic calcium led to CaM KKβ–dependent activation of AMPK to improve glucose metabolism. However, altered calcium handling prevented normal muscle relaxation, leading to disruption of subsequent contraction manifested as decreased overall muscle performance. At the sarcomere, cytosolic calcium binds troponin to release tropomyosin from actin, thus permitting myosin-induced ATP-dependent muscle contraction. During normal relaxation, cytosolic calcium is taken up by the SR, leading to actin-myosin dissociation. Calcium cycling between SR and the cytosol is strictly regulated in striated muscles, and its disruption causes muscle dysfunction (43). Figure 12 depicts the potential relationship among FAS, insulin sensitivity, and weakness in skeletal muscle.

FAS is linked to the generation of phospholipid species that activate nuclear receptors (17, 44), consistent with the possibility that FAS is part of a synthetic pathway leading to the production of phospholipids, such as certain PE species, at the SR. However, our data did not establish that FAS enzyme activity is required for effects on SR composition. It is possible that FAS interacts with the SR or SERCA to alter the PC/PE ratio independent of its capacity to convert malonyl-CoA into palmitate.

Lipids are implicated in the pathogenesis of diet-induced skeletal muscle insulin resistance. This phenomenon is incompletely understood, but a popular explanation is the accumulation of lipid species such as DAG and ceramide that may induce stress kinases, leading to decreased insulin-stimulated Akt phosphorylation (24, 45). Perhaps underappreciated is reduced muscle AMPK activity in insulin resistance (29, 30). Skeletal muscles from FASKOS mice were not protected from HFD-induced suppression of insulin-stimulated Akt phosphorylation, but they were protected from a HFD-induced reduction in AMPK activity. DAG, ceramide, and the activities of certain stress kinases were not different in muscles from HFD-fed control and FASKOS mice, which suggests that these mediators are not likely contributors to the insulin sensitivity of FASKOS muscle. Our observations indicate that the insulin-sensitizing phenotype with FAS deficiency is mediated by AMPK, not Akt, and suggest that multiple pathways may be involved in muscle insulin resistance.

FAS affects activity of the nuclear receptor PPARα in liver (17, 22), brain (46), and macrophages (47), and overexpression of PPARα in skeletal muscle causes insulin resistance and inactivates AMPK (18), which led us to consider PPARα as a mediator of the FASKOS phenotype. However, FAS inactivation in muscle did not affect activity of a PPARα reporter gene, and treatment
with a potent PPARα agonist did not rescue the effect of FAS deletion on AMPK signaling. Similarly, inactivation of FAS in cardiac muscle did not affect PPARα target genes, but instead activated calcium signaling pathways (33) through incompletely defined mechanisms. The calcium-regulatory mechanism shown here in skeletal muscle may also be involved in the phenotype of mice with FAS knockout in heart (33), another tissue adapted for contractile function, but its relevance to other tissue-specific FAS knockouts is uncertain. Inactivation of pancreatic β cell FAS has no effect on insulin secretion, a process requiring dynamic

**Figure 7**

Muscle FAS deletion does not affect PPARα activity, but increases cytosolic calcium. (A) Western blot analyses of scrambled (SC) or FASKD C2C12 myocytes. (B) PPARα luciferase activity in scrambled or FASKD C2C12 myocytes with or without the PPARα agonist WY14643 (n = 8 replicate measurements per bar). (C) Western blot analyses of scrambled or FASKD C2C12 myocytes with or without WY14643. (D) Cytosolic calcium in scrambled and FASKD C2C12 myocytes measured using fluo-8 AM. Original magnification, ×100. (E) Quantified fluo-8 AM fluorescence in scrambled and FASKD C2C12 myocytes (n = 12 culture wells per condition; horizontal lines denote means). Data are mean ± SEM.

**Figure 8**

Muscle FAS deletion induces calcium signaling and activates AMPK through CaMKKβ. (A) Calcium signaling cascade affected by muscle FAS deletion. PLB, phospholamban. (B–D) Western blot analyses for calcium signaling proteins (n = 6 per group). Data are mean ± SEM. Representative blots are also shown. (B) C2C12 myocytes. Phospholamban is phosphorylated by CaMKII on Thr17 and by PKA on Ser16. (C) Muscle from clamp studies of HFD-fed animals. (D) Isolated soleus muscle. (E) Western blot analyses of C2C12 myocytes after FASKD and CaMKKβ knockdown. (F) Incubation of C2C12 myocytes with the CaMKK inhibitor STO-609 (10 μg/ml for 6 hours) eliminated FASKD-induced activation of AMPK signaling.
Figure 9
Muscle FAS deletion decreases SERCA activity. (A) Differential centrifugation of mouse hindlimb muscles. PM, plasma membrane; GM130, 130-kDa cis-Golgi matrix protein. (B) Soleus co-IP experiments. FAS, but not ACC, coimmunoprecipitated with SERCA1 or SERCA2 antibody; SERCA1 and SERCA2 coimmunoprecipitated with FAS antibody, but not with ACC antibody. (C) Confocal images of C2C12 cells. Original magnification, ×630. Boxed regions are shown enlarged (×4-fold) at far right. (D) Rescue of the FASKD-induced reduction in SERCA enzyme activity by expression of myc-SERCA1 (n = 8 replicates per condition). Moreover, SR SERCA abundance with vector treatment was not altered by FASKD (bottom). (E) Expression of myc-SERCA1 reversed FASKD-induced increase in pAMPK. (F) SERCA activity in soleus from chow- or HFD-fed control and FASKOS mice (n = 6 per group). SR SERCA abundance is also shown. Data are mean ± SEM.
regulation of intracellular calcium (46), which suggests that in this model, the loss of FAS does not have a major impact on the calcium flux required to release insulin-secretory granules. However, it is possible that the loss of FAS in other nonmuscle tissues alters calcium signaling.

Our data suggest that the induction of FAS by HFD feeding may contribute to skeletal muscle insulin resistance. Although not extensively studied in muscle, FAS regulation involves insulin signaling, the induction of the transcription factor SREBP1c (also known as SREBF1), and other factors (48). In humans, skeletal muscle expression of SREBP1c is highest in individuals that are most insulin resistant (49). This observation is consistent with our present findings in mice, supporting the plausibility of SREBP1c-driven genes such as FAS being increased in muscle in the setting of diet-induced insulin resistance. HFD feeding has been reported to increase SERCA activity in soleus muscle under certain conditions (50), also consistent with our observations linking HFD feeding, FAS, and SERCA.

It is tempting to speculate that HFD-induced FAS expression in muscle represents a selective advantage for mammals. Skeletal muscle’s raison d’être is contraction, required for acquiring food, escaping predators, and finding a mate. A HFD-induced increase in muscle FAS may promote insulin resistance, but it may also optimize contractile function. When evolutionary pressures were substantial, mammalian meals were infrequent but high in fat, especially if they consisted of other mammals. Our ancestors probably fasted for prolonged periods while hunting, then feasted for prolonged periods on high-fat prey. Disruption of metabolic homeostasis induced by HFD feeding after fasting may have prompted the selection of animals with the capacity to increase FAS activity in muscle and maintain SERCA activity in order to escape other predators attracted by the same high-fat prey. It has been suggested that insulin resistance is a normal physiological response that limits obesity through mobilization of substrates (51). Our results suggest that, in addition to defending against obesity, insulin resistance maintains muscle strength in the setting of nutrient excess. A HFD-induced increase in skeletal muscle FAS would preserve SERCA activity and normalize force-generating capacity, perhaps selected by evolutionary pressures. It would also increase insulin resistance, probably unrelated to evolutionary pressures since the complications of insulin resistance become manifest well after the next generation is spawned.

Diabetes may have complex effects on skeletal muscle calcium metabolism, since SERCA is decreased in the diabetic soleus muscle (52) and insulin receptor substrate proteins interact with SERCA (53). FAS inhibitors may be useful for treating type 2 diabetes (54). Our findings suggest that these agents might lower blood sugar at the expense of muscle function, identifying a possible side effect of a new class of medication, but also representing a potentially reasonable exchange for modern humans who need to escape the consequences of abnormal glucose metabolism instead of another mammal looking for a meal.

**Methods**

Further details are provided in Supplemental Methods.

**Animals.** Mice with a floxed Fasn locus (Fasn<sup>fl/fl</sup>) were crossed with transgenic mice expressing Cre recombinase under the control of the human α-skeletal actin (HSA) promoter to obtain FASKOS mice in the C57BL/6J strain.

**Figure 10**

Muscle FAS deletion alters SR phospholipid composition. Mass spectrometry quantification of (A) relative PE and PC abundance, (B and C) total PE and PC, and (D and E) PE and PC species in isolated SR from hindlimb muscles of chow- or HFD-fed control and FASKOS mice (n = 4–6). Data are mean ± SEM.
cDNA insert (Open Biosystems) by PCR, using the following primers (underline denotes XhoI and XbaI sites): 5′-CTCGAGATGGAGGCCGCGCACTC-CAAGTCC-3′ and 5′-TCTAGATTATCCCTCCAGATAGTTCCGAGC-3′. The amplified product was subcloned into the pRK5-myc plasmid at its XhoI (5′) and XbaI (3′) sites to yield an N-terminal myc-tagged construct (pRK5-myc-mSERCA1). Transfection-ready pCMV6-XL4-SERCA2 plasmids were purchased from OriGene Technologies. All plasmids were sequenced to confirm their identity and orientation. Differentiated scrambled or FASKD C2C12 myocytes were transfected with pRK5-myc-mSERCA1 or pCMV6-XL4-SERCA2 using Lipofectamine 2000 (Invitrogen).

**IP.** Homogenates were incubated with antibodies (anti-FAS [ab22759], anti-SERCA1 [ab2818], and anti-SERCA2 [ab3625]; Abcam) or normal rabbit serum on a rotator overnight at 4°C. Protein A agarose beads were added to the homogenate-antibody mixture for an additional 2 hours at 4°C. Beads were washed 5 times with PBS, mixed with 2× Laemmli sample buffer, heated at 95°C for 5 minutes, and then subjected to SDS-PAGE for analysis by Western blotting or mass spectrometry.

**SERCA activity assay.** The calcium transport assay for measuring SERCA activity was adapted from a protocol described previously (58). SR fractions from C2C12 myocytes or skeletal muscles were isolated by differential centrifugation. The assay mixture contained 100 mM KCl, 30 mM imidazole-histidine (pH 6.8), 5 mM MgCl2, 5 mM ATP, 5 mM (COOK)2, 5 mM NaN3, 1 μM Rethenium Red, and 50 μM CaCl2 (1 μCi/μmol [45Ca]CaCl2; American Radiolabeled Chemicals). The reaction was started by the addition of SR fractions containing 150 μg protein for 15 minutes in a 37°C water bath, stopped by the addition of 0.15 M KCl and 1 mM LaCl3, and filtered through a 0.2-μm HT Tuffryn membrane ( Pall Corp.). SERCA-independent calcium transport was quantified in the presence of 10 μM thapsigargin.

**Indirect calorimetry.** Indirect calorimetry was performed as described previously (55). O2 consumption (VO2) and CO2 production were measured for 24 hours. Mean relative VO2 (expressed in ml/kg/min) and respiratory quotient (RQ; calculated as VCO2/VO2) were determined for dark and light periods.

**AMP and ATP measurements.** Frozen muscles were treated with perchloric acid and neutralized with K2CO3 on ice, and then levels were assayed by HPLC as described previously (56). AMP and ATP were quantified based on peak area compared with a standard curve and normalized to frozen tissue weight.

**Fatty acid oxidation determinations.** Muscles or cells were processed exactly as described previously (44) in an assay after the conversion of labeled palmitate to labeled CO2 that is trapped in filter paper. Rates of fatty acid oxidation were calculated as mmol 13C-labeled CO2 produced per hour, normalized to tissue weight or cellular protein.

**Treadmill sprint interval exercise tests.** Mouse treadmill exercise tests used a recently described protocol (57). Fed mice ran intervals of 1 minute each, with a 2-minute rest between each interval. Running started at a rate of 10 m/min and increased by 5 m/min at each interval.

**Ectopic expression of SERCA1 and SERCA2.** The cDNA encoding the mouse enzyme SERCA1 was cloned from pCMV-SPORT6 with mouse SERCA1 cDNA insert (Open Biosystems) by PCR, using the following primers (underline denotes XhoI and XbaI sites): 5′-CTCGAGATGGAGGCCGCGCACTC-CAAGTCC-3′ and 5′-CTTAGATTATCCCTCCAGATAGTTCCGAGC-3′. The amplified product was subcloned into the pRK5-myc plasmid at its XhoI (5′) and XbaI (3′) sites to yield an N-terminal myc-tagged construct (pRK5-myc-mSERCA1). Transfection-ready pCMV6-XL4-SERCA2 plasmids were purchased from OriGene Technologies. All plasmids were sequenced to confirm their identity and orientation. Differentiated scrambled or FASKD C2C12 myocytes were transfected with pRK5-myc-mSERCA1 or pCMV6-XL4-SERCA2 using Lipofectamine 2000 (Invitrogen).

**FASN and Cre genotyping was performed as described previously (22). Chow diet and HFD were Purina 4043 and Harlan Teklad TD 88137, respectively. Male mice aged 8–12 weeks were used for all experiments. Controls were either Fasnfl/fl mice without the Cre transgene or mice with the Cre transgene alone. The presence of Cre transgene alone did not affect glucose metabolism compared with Fasnfl/fl mice without the transgene (data not shown).**

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**Ectopic expression of SERCA1 and SERCA2.** The cDNA encoding the mouse enzyme SERCA1 was cloned from pCMV-SPORT6 with mouse SERCA1 cDNA insert (Open Biosystems) by PCR, using the following primers (underline denotes XhoI and XbaI sites): 5′-CTCGAGATGGAGGCCGCGCACTC-CAAGTCC-3′ and 5′-CTTAGATTATCCCTCCAGATAGTTCCGAGC-3′. The amplified product was subcloned into the pRK5-myc plasmid at its XhoI (5′) and XbaI (3′) sites to yield an N-terminal myc-tagged construct (pRK5-myc-mSERCA1). Transfection-ready pCMV6-XL4-SERCA2 plasmids were purchased from OriGene Technologies. All plasmids were sequenced to confirm their identity and orientation. Differentiated scrambled or FASKD C2C12 myocytes were transfected with pRK5-myc-mSERCA1 or pCMV6-XL4-SERCA2 using Lipofectamine 2000 (Invitrogen).

**IP.** Homogenates were incubated with antibodies (anti-FAS [ab22759], anti-SERCA1 [ab2818], and anti-SERCA2 [ab3625]; Abcam) or normal rabbit serum on a rotator overnight at 4°C. Protein A agarose beads were added to the homogenate-antibody mixture for an additional 2 hours at 4°C. Beads were washed 5 times with PBS, mixed with 2× Laemmli sample buffer, heated at 95°C for 5 minutes, and then subjected to SDS-PAGE for analysis by Western blotting or mass spectrometry.

**SERCA activity assay.** The calcium transport assay for measuring SERCA activity was adapted from a protocol described previously (58). SR fractions from C2C12 myocytes or skeletal muscles were isolated by differential centrifugation. The assay mixture contained 100 mM KCl, 30 mM imidazole-histidine (pH 6.8), 5 mM MgCl2, 5 mM ATP, 5 mM (COOK)2, 5 mM NaN3, 1 μM Rethenium Red, and 50 μM CaCl2 (1 μCi/μmol [45Ca]CaCl2; American Radiolabeled Chemicals). The reaction was started by the addition of SR fractions containing 150 μg protein for 15 minutes in a 37°C water bath, stopped by the addition of 0.15 M KCl and 1 mM LaCl3, and filtered through a 0.2-μm HT Tuffryn membrane ( Pall Corp.). SERCA-independent calcium transport was quantified in the presence of 10 μM thapsigargin.

**Indirect calorimetry.** Indirect calorimetry was performed as described previously (55). O2 consumption (VO2) and CO2 production were measured for 24 hours. Mean relative VO2 (expressed in ml/kg/min) and respiratory quotient (RQ; calculated as VCO2/VO2) were determined for dark and light periods.

**AMP and ATP measurements.** Frozen muscles were treated with perchloric acid and neutralized with K2CO3 on ice, and then levels were assayed by HPLC as described previously (56). AMP and ATP were quantified based on peak area compared with a standard curve and normalized to frozen tissue weight.

**Fatty acid oxidation determinations.** Muscles or cells were processed exactly as described previously (44) in an assay after the conversion of labeled palmitate to labeled CO2 that is trapped in filter paper. Rates of fatty acid oxidation were calculated as mmol 13C-labeled CO2 produced per hour, normalized to tissue weight or cellular protein.

**Treadmill sprint interval exercise tests.** Mouse treadmill exercise tests used a recently described protocol (57). Fed mice ran intervals of 1 minute each, with a 2-minute rest between each interval. Running started at a rate of 10 m/min and increased by 5 m/min at each interval.

**Ectopic expression of SERCA1 and SERCA2.** The cDNA encoding the mouse enzyme SERCA1 was cloned from pCMV-SPORT6 with mouse SERCA1
FAS activity assay. FAS enzyme activity was determined using a modification of a previously described assay (59). Freshly harvested muscles were weighed and homogenized in 0.1 M potassium phosphate buffer (pH 7.0) containing 8% sucrose, 1 mM EDTA (pH 8.0), and 20 mM 2-mercapto-ethanol at 4°C. Homogenates were centrifuged at 3,000 g for 10 minutes at 4°C. 10 µl of the supernatant was added to 80 µl of assay buffer (0.14 M potassium phosphate buffer [pH 7.0], 1.4 mM EDTA [pH 8.0], 1.4 mM DTT, 0.24 mM NADPH, and 0.1 mM acetyl-CoA), and the rate of NADPH oxidation was monitored at 340 nm. The substrate-dependent rate was determined by subtracting the NADPH oxidation rate from the rate after adding 10 µl of 1 mM malonyl-CoA.

Measurement of malonyl-CoA content. Soleus muscles were harvested from mice after a 6-hour fast. Malonyl-CoA was determined in extracts homogenized in deionized water. CoA esters were separated using reverse-phase HPLC on a 5 µm Supelco C18 column with a Waters HPLC system, monitoring 254 nm as the maximal absorbance for CoA. Buffers, gradients, and flow rates were described previously (60).

Statistics. Values are expressed as mean ± SEM. Statistical comparisons were performed using an unpaired 2-tailed Student’s t test (for 2-group analyses) or 2-way ANOVA with Student-Newman-Keuls post-hoc test (for 2-by-2 comparisons). A P value less than 0.05 was considered significant.

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