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Ceramide synthase 5 mediates lipid-induced autophagy and hypertrophy in cardiomyocytes

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Diabetic cardiomyopathy (DbCM), which consists of cardiac hypertrophy and failure in the absence of traditional risk factors, is a major contributor to increased heart failure risk in type 2 diabetes patients. In rodent models of DbCM, cardiac hypertrophy and dysfunction have been shown to depend upon saturated fatty acid (SFA) oversupply and de novo sphingolipid synthesis. However, it is not known whether these effects are mediated by bulk SFAs and sphingolipids or by individual lipid species. In this report, we demonstrate that a diet high in SFA induced cardiac hypertrophy, left ventricular systolic and diastolic dysfunction, and autophagy in mice. Furthermore, treatment with the SFA myristate, but not palmitate, induced hypertrophy and autophagy in adult primary cardiomyocytes. De novo sphingolipid synthesis was required for induction of all pathological features observed both in vitro and in vivo, and autophagy was required for induction of hypertrophy in vitro. Finally, we implicated a specific ceramide N-acyl chain length in this process and demonstrated a requirement for (dihydro)ceramide synthase 5 in cardiomyocyte autophagy and myristate-mediated hypertrophy. Thus, this report reveals a requirement for a specific sphingolipid metabolic route and dietary SFAs in the molecular pathogenesis of lipotoxic cardiomyopathy and hypertrophy.

Introduction
Obesity and diabetes present two of the most important health challenges facing the Western world at this time. Patients suffering from type 2 diabetes (T2D) are subject to a number of major health risks, including a greatly increased risk of heart failure (1). This is due in part to the development of diabetic cardiomyopathy (DbCM), which occurs independently of other traditional risk factors (2). DbCM promotes cardiac remodeling and impairs cardiac function (1). Importantly, individuals with T2D and the metabolic syndrome present with dyslipidemia, and recent studies have suggested that DbCM may occur as a result of lipid overload and subsequent lipotoxic events (ref. 2, reviewed in ref. 3). In particular, oversupply of saturated fatty acids (SFAs) has been implicated in this process.

Previous studies of lipotoxic DbCM in rodents have relied on several important transgenic models. The B6.Cg-Lep°/° (ob/ob) and B6.BKS(D)-Leprdb/db (db/db) mouse models, which lack the genes encoding leptin and the leptin receptor, respectively, are both very popular models that develop obesity and a DbCM-like cardiac phenotype (reviewed in ref. 4). Other less common models, including the Atg7-knockout mouse and the LpL−/− transgenic mouse, also induce lipotoxic cardiomyopathy by perturbing cardiac lipid uptake, handling, or metabolism (reviewed in ref. 4). While these transgenic models robustly induce lipid overload in cardiomyocytes, the very disruptions that produce lipotoxic DbCM also drastically alter patterns of lipid uptake and handling in a nonphysiologic way. In contrast, wild-type mice fed standard lard-based high-fat diets (LBD) failed to develop a DbCM-like phenotype until very late time points, if at all, and insulin resistance and other aspects of the diabetic phenotype were less pronounced in this model system than in transgenic animals (4–9). This may result from the high levels of protective unsaturated fatty acids (UFAs) contained in these diets (10–12). Thus, while these transgenic model systems produce a robust DbCM phenotype, they are unable to reveal clinically relevant roles of specific lipid species in the molecular pathogenesis of DbCM.

Because of these weaknesses in present model systems, our laboratory recently implemented a new obesogenic high-fat diet that was designed to replicate the shift toward shorter SFAs observed in the plasma of obese and diabetic patients (13). Milk was considered an ideal fat source for this diet, as it contains low levels of UFAs and high levels of saturated fat. Furthermore, epidemiologic studies have demonstrated a relationship between consumption of high-fat dairy (e.g., butter, cheese, ice cream) and insulin resistance in humans (reviewed in ref. 14). Similarly, multiple studies have also linked the SFA myristate (C14:0), which is a major constituent of milk fat, to insulin resistance and the metabolic syndrome in humans (14–17). Strikingly, in wild-type C57BL/6J mice, this milk fat–based diet (MFBD) promoted profound insulin resistance and hyperglycemia by 8 weeks. This stands in contrast with mice fed a traditional LBD, which do not develop metabolic defects until very late time points (4). These findings suggested that the MFBD might promote a more severe DbCM phenotype than the LBD in wild-type mice.

Additional evidence suggests that milk fat and its constituent fatty acid, myristate, may promote cardiac dysfunction and hypertrophy. In particular, dietary patterns that feature high-fat dairy were associated with increased LV mass and reduced systolic function in humans, and plasma myristate content was positively associated with incidence of heart failure and total cardiovascular disease mortality (18–20). Furthermore, a recent study demonstrated that infusion with a myristate-rich SFA cocktail rapidly induced LV hypertrophy in mice (21). These data suggest a role for dietary and plasma myristate in cardiac hypertrophy and dysfunction, but potential mechanisms remain unknown.

Conflict of interest: The authors have declared that no conflict of interest exists.
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Thus, the literature has provided epidemiological links between dietary milk fat, the fatty acid myristate, and both cardiac dysfunction and diabetes in humans, while data in animals suggest a potential role for myristate in cardiac hypertrophy. Based on these findings, we hypothesized that the MFBD might induce a cardiac phenotype closely resembling human DbCM, with no requirement for genetic perturbations. Furthermore, we hypothesized that this phenotype would develop in a sphingolipid-dependent manner.

This report characterizes the cardiac phenotype of mice fed the MFBD, which is rich in SFAs but not UFAs, and implicates a specific SFA and sphingolipid metabolic route in the induction of hypertrophy and increased autophagy in response to lipid overload in cardiomyocytes.

Results
An MFBD induces cardiac hypertrophy and dysfunction. Based on the documented connection between diets high in saturated fat and both insulin resistance and incidence of heart disease in humans, we hypothesized that the MFBD, which is high in SFAs and low in protective UFAs, would induce DbCM-like cardiac hypertrophy and dysfunction in wild-type mice. To test this hypothesis, mice were maintained on the MFBD, a traditional LBD, or an isocaloric low-fat control diet (CD) for fifteen weeks and then subjected to echocardiographic analyses. By 15 weeks of high-fat feeding, MFBD mice developed LV hypertrophy and functionally significant reductions in ejection fraction relative to mice fed the CD (Table 1). In contrast, LBD-fed mice maintained normal cardiac function and LV mass relative to CD-fed mice. These results suggested that a diet rich in saturated fat but not unsaturated fat promoted LV hypertrophy and diminished cardiac function.

These findings led us to ask whether these observations constituted a pure DbCM phenotype, i.e., one independent of pressure overload and cardiac dilation. To determine this, blood pressure measurements, detailed echocardiographic studies, and histological assessments were performed on these mice after 18 weeks of high-fat feeding. Despite similar BWs, blood pressures, and end-diastolic diameters between high-fat diet groups (Figure 1A and Table 2), only MFBD mice displayed reduced ejection fraction, reduced early diastolic mitral annular velocity, increased end-diastolic posterior wall thickness, and LV hypertrophy (Table 2). Furthermore, only mice fed the MFBD displayed whole heart, LV, and RV hypertrophy and increased cardiomyocyte cross-sectional area (Figure 1, B and C). These findings indicate that the MFBD, but not the LBD or CD, induced concentric cardiac hypertrophy and reduced cardiac function in the absence of dilated cardiomyopathy or pressure overload.

Cardiac outcomes in the MFBD are sphingolipid dependent. Mechanistically, lipotoxicity has been implicated in the pathogenesis of DbCM (22). Previous studies in mice overexpressing a membrane-anchored lipoprotein lipase in the heart indicated that cardiac dysfunction, hypertrophy, and expression of heart failure markers were ameliorated by inhibition of de novo sphingolipid synthesis; however, the clinical relevance of this model to human DbCM may be limited (23). Moreover, only bulk sphingolipid synthesis has been implicated in these effects; the specific molecular species of sphingolipids and the enzymes that mediate them have remained undetermined.

Table 1
Echocardiographic assessment of mice after 15 weeks of high-fat diet

<table>
<thead>
<tr>
<th>Diet</th>
<th>BW (g)</th>
<th>EF (%)</th>
<th>LV mass (g)</th>
<th>LV mass/Tibia length (g/mm)</th>
<th>ED PWTH (mm)</th>
<th>RWTH</th>
<th>EDD (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD</td>
<td>32.1 ± 1.4</td>
<td>65 ± 1</td>
<td>85.7 ± 2.8</td>
<td>4.3 ± 0.1</td>
<td>0.74 ± 0.01</td>
<td>0.37 ± 0.01</td>
<td>4.0 ± 0.1</td>
</tr>
<tr>
<td>MFBD</td>
<td>38.7 ± 2.2</td>
<td>54 ± 1A</td>
<td>102.4 ± 4.0A</td>
<td>5.1 ± 0.2A</td>
<td>0.88 ± 0.02A</td>
<td>0.45 ± 0.01</td>
<td>3.9 ± 0.1</td>
</tr>
<tr>
<td>LBD</td>
<td>37.8 ± 2.7</td>
<td>62 ± 1</td>
<td>79.8 ± 5.4</td>
<td>4.0 ± 0.3</td>
<td>0.74 ± 0.03</td>
<td>0.39 ± 0.02</td>
<td>3.8 ± 0.1</td>
</tr>
</tbody>
</table>

Echocardiographic measurements demonstrated reduced ejection fraction and increased LV mass in mice fed an MFBD, which is high in saturated fat. EDD, end-diastolic dimension; EF, ejection fraction; TL, tibia length; ED PWTH, end-diastolic posterior wall thickness. Data are presented as mean ± SEM, A*P < 0.05 vs. control.

![Figure 1](image-url)

Figure 1
An MFBD promotes sphingolipid-dependent cardiac hypertrophy. Mice fed an MFBD for 18 weeks (A) did not experience greater obesity than mice on a traditional LBD, but they displayed (B) marked cardiac hypertrophy and (C) increased cardiomyocyte cross-sectional area. Myriocin treatment attenuated this hypertrophy without affecting weight gain. Black bars, control plus DMSO; light gray bars, milk fat plus DMSO; dark gray bars, milk fat plus myriocin; hatched bars, lard plus DMSO. All results are given as mean ± SEM. *P < 0.05 vs. control.
To determine whether the cardiac remodeling observed in MFBD mice was dependent on sphingolipid synthesis, MFBD mice were treated with the de novo sphingolipid synthesis inhibitor myriocin and subjected to echocardiography and histological assessment. Strikingly, inhibition of de novo sphingolipid synthesis restored all measured echocardiographic parameters to normal levels in the MFBD mice (Table 2). Furthermore, myriocin treatment prevented LV and cardiomyocyte hypertrophy without reducing BW (Figure 1). These results dramatically highlight a role for sphingolipids in the pathogenesis of concentric cardiac hypertrophy and cardiac dysfunction in diet-induced obesity and DbCM.

Because attenuation of de novo sphingolipid synthesis prevented development of cardiomyopathy in the MFBD, we hypothesized that the cardiomyopathy phenotype may result from increases in bulk ceramide levels. To test this, ceramide and its precursor dihydroceramide were measured in hearts of mice fed the MFBD or the LBD. Surprisingly, total ceramide and dihydroceramide did not increase in the MFBD, but rather increased in the less cardiotoxic LBD, relative to control (Figure 2A). This perplexing result suggested a more subtle mechanism for sphingolipid-dependent lipotoxicity than merely a simple increase in bulk ceramide levels, as had previously been assumed.

The term “ceramide” generally refers to any simple N-acylated sphingosine moiety; however, many different N-acyl chain lengths occur in ceramides, with the natural range extending from 12 to 26 carbons (24). Importantly, recent studies indicate that elevations of specific ceramide N-acyl chain lengths promote particular cellular and signaling outcomes in some systems (25, 26). In order to determine whether the MFBD induced changes in specific ceramides, individual ceramide species were quantified in hearts from mice after 8 weeks of high-fat feeding. This time point was chosen to minimize confounding effects of long-term insulin resistance, obesity, and inflammation. Strikingly, the MFBD specifically increased C14-ceramide (Figure 2B); similar results were obtained at later time points (data not shown). This finding was consistent with the lipid content of the MFBD, which contained significantly more myristate than either the CD or the LBD and promoted a 4-fold increase in plasma myristate levels (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI63888DS1). These results indicate that the MFBD induced a very precise readjustment of ceramide N-acyl chain length composition, possibly consequent to elevated plasma myristate levels.

Myristate induces hypertrophy through ceramide synthase 5. Because of the high myristate content of the MFBD, we were led to ask whether exposure to myristate alone might induce hypertrophy in cardiomyocytes. To test this, we measured the effects of myristate treatment on cell size in isolated primary cardiomyocytes. Consistent with a specific role for myristate in cardiac hypertrophy, myristate treatment increased cell size in isolated cardiomyocytes (Figure 3A). Furthermore, as in the MFBD hearts, this hypertrophic response was completely attenuated by inhibition of de novo sphingolipid synthesis with myriocin. In contrast, treatment with palmitate, a fatty acid implicated in lipotoxicity

![Figure 2](image-url)

An MFBD production of C14-ceramide. (A) Total ceramide (Cer) and dihydroceramide (DHC) did not increase in mice fed an MFBD for 8 weeks, but (B) C14-ceramide increased specifically in the MFBD but not the LBD. All results are given as mean ± SEM. *P < 0.05 vs. control.
in other tissues, did not induce cardiomyocyte hypertrophy. These findings suggest a specific role for myristate or its derivatives in the induction of sphingolipid-dependent cardiomyocyte hypertrophy.

The specific upregulation of C14-ceramide in the MFBD myoccardium suggested that this metabolite, or the pathway that synthesizes it, may play a unique role in lipotoxic cardiac hypertrophy. Supporting this hypothesis, myristate treatment, which promoted cardiomyocyte hypertrophy, increased C14-ceramide levels 10-fold in isolated cardiomyocytes (Figure 3B). In contrast, palmitate treatment, which did not induce hypertrophy, had no effect on C14-ceramide levels. These findings led us to ask whether the pathway that produces C14-ceramide might be required for myristate-induced cardiomyocyte hypertrophy.

Ceramide synthesis occurs through N-acylation of a sphingoid base by one of 6 ceramide synthase (CerS) isoforms, each of which displays a unique profile of N-acyl chain length incorporation (Figure 3C) (27, 28). While both CerS5 and CerS6 synthesize C14-ceramide robustly in vitro (28), expression of CerS6 is very low in cardiomyocytes. Thus, these findings implicate a specific CerS isoform in a sphingolipid-dependent mechanism.

Figure 3
Myristate induces hypertrophy in a sphingolipid- and CerS5-dependent manner. (A) Myristate, but not palmitate, treatment induced hypertrophy in cardiomyocytes. This was prevented by inhibition of de novo sphingolipid synthesis with myriocin. Results are shown as percentage increase over BSA. (B) Myristate, but not palmitate, strongly induces C14-ceramide production in cardiomyocytes. (C) CerS5 and CerS6 are primarily responsible for synthesis of C14-ceramide. (D) siRNA-mediated knockdown of CerS5 is sufficient to prevent induction of hypertrophy by myristate. Results are given as percentage increase over control siRNA treated with BSA. For all panels, results are presented as mean ± SEM. *P < 0.05 vs. BSA; **P < 0.005 vs. control BSA; ***P < 0.05 vs. palmitate; †P < 0.005 vs. myristate DMSO.

The MFBD induces autophagy in a sphingolipid-dependent manner. Numerous studies have identified autophagy as a key process that is upregulated in cardiac hypertrophy in response to pressure overload (31–36). Furthermore, sphingolipid-dependent autophagy has been demonstrated in numerous cell types, with the molecular mechanisms appearing to be specific to both the cell type and the mode of induction (reviewed in refs. 37–39). However, few studies have addressed the role of autophagy in DbCM in T2D, and none of them have directly tested the effect of lipid oversupply on cardiac autophagy.

To determine whether the MFBD increased autophagy, expression of the autophagy markers was measured in whole-heart lysate at 8 weeks. Indeed, mRNA levels of LC3B (Map1lc3b) and Beclin 1 (Becn1) were both strongly upregulated in the MFBD (Figure 4A). Furthermore, immunohistochemical and immunofluorescent staining demonstrated increased numbers of LC3B puncta in the LV of mice fed the MFBD, but not the LBD (Figure 4, B and C); moreover, this effect was attenuated in myriocin-treated mice, suggesting that autophagy in the MFBD mice occurred through a sphingolipid-dependent mechanism.

Myristate induces autophagy through ceramide synthase 5. Because of the high myristate content of the MFBD and the confirmed role of exogenous myristate in the induction of cardiomyocyte hypertrophy in this system, we were led to ask whether exposure to myristate might also induce autophagy in cardiomyocytes. To test this, we treated isolated adult primary cardiomyocytes with myristate and measured levels of autophagy markers. Consistent with observations in whole heart, myristate treatment promoted overexpression of Becn1 and Atg7, another marker of autophagy, compared with BSA (Figure 5, A and B). In contrast, treatment with palmitate (a C16:0 fatty acid) did not induce expression of autophagy markers. These findings strongly paralleled those regarding the role of myristate in cardiomyocyte hypertrophy. Additionally, a prior study indicated that C14-ceramide and CerS5 increased under proautophagic conditions in cancer cells, although the potential mechanistic link was not investigated (40).
Thus, we were led to ask whether CerS5 might also mediate induction of cardiomyocyte autophagy by myristate.

Previous studies have indicated that upregulation of BECN1 expression was sufficient to increase cardiac autophagy, both basally and in response to stress, while diminishing BECN1 expression attenuated stress-induced autophagy (34). Consistent with a role for CerS5 in regulation of cardiomyocyte autophagy in the present system, overexpression of CerS5 induced BECN1 expression in isolated cardiomyocytes, even in the absence of myristate treatment (Figure 5C). Furthermore, siRNA-mediated knockdown of endogenous CerS5 reduced basal expression of BECN1 (Figure 5D). To test directly the potential role of CerS5 in sphingolipid-mediated cardiomyocyte autophagy, we treated cells with the sphingoid base dihydrosphingosine (DHS), which serves as the immediate precursor of dihydroceramide. Because all CerS isoforms utilize DHS as a substrate, CerS5 knockdown would only prevent autophagy if that isoform were specifically required for this process. Supporting a role for a CerS-mediated sphingolipid pathway in cardiac autophagy, treatment with DHS induced BECN1 expression in isolated cardiomyocytes (Figure 5D). Strikingly, CerS5 knockdown completely prevented induction of BECN1 by DHS. This explicitly confirmed the requirement for this CerS isoform in sphingolipid-induced BECN1 expression in cardiomyocytes. Finally, to confirm the role for CerS5 in myristate-induced autophagy per se, myristate-treated cells were subjected to siRNA-mediated CerS5 knockdown. Consistent with the preceding results, CerS5 knockdown abrogated the induction of BECN1 expression by myristate treatment (Figure 5E). These findings demonstrated a requirement for CerS5 in sphingolipid-mediated induction of Beclin 1 protein and message levels in cardiomyocytes.

As suggestive as these results might be, they do not definitively demonstrate that myristate induces autophagy in cardiomyocytes. Indeed, it is possible that levels of autophagy mediators might be upregulated due to a defect in fusion of the autophagosome to the lysosome rather than increased autophagic flux. To test this possibility, autophagic flux was assessed in 2 complementary approaches, both of which utilized a GFP-LC3B fusion construct.
The first approach used the GFP-LC3B fusion construct to track LC3B lipidation and breakdown. The GFP-LC3B fusion protein is lipidated and processed similarly to native LC3B (41); however, the GFP moiety is more resistant to lysosomal proteases than the LC3B moiety, causing the free GFP protein to persist in the lysosome after breakdown of LC3B. Thus, levels of free GFP, detected by Western blot, can be used as a readout for breakdown of the inner membrane and contents of the autophagosome (41). Similarly to the results obtained with BECN1, it was found that overexpression of CerS5 in isolated cardiomyocytes resulted in an increase in Beclin 1 mRNA and protein (Figure 6A). Overexpression of CerS5 in isolated cardiomyocytes increased expression of GFP-LC3B-I 4-fold, while GFP-LC3B-II and free GFP expression increased 2- and 3-fold, respectively. Additionally, treatment with myristate increased levels of GFP-LC3B-II 7-fold and levels of free GFP 13-fold compared with BSA controls (Figure 6B). Furthermore, it was revealed that total GFP-labeled puncta increased in myristate-treated cells, compared with those treated with BSA, and this increase was attenuated by CerS5 knockdown (Figure 6C). Taken together, all of these results demonstrate a requirement for CerS5 in induction of sphingolipid-mediated autophagic flux by myristate.

In a complementary approach, autophagosome dynamics were examined by fluorescent microscopy. To track autophagosome formation and fusion with the lysosome, GFP-LC3B–expressing cells were treated with myristate or BSA for 16 hours, loaded with the acidotropic dye LysoTracker Red, and examined by fluorescence microscopy over a 20-minute time course. This approach did not provide evidence of aberrant autophagosome accumulation and, rather, confirmed that fusion of autophagosomes to the lysosome, as indicated by fluorescent colocalization, is unimpaired in myristate-treated cells (Figure 7 and Supplemental Figures 5 and 6). Furthermore, it was revealed that total GFP-labeled puncta increased in myristate-treated cells, compared with those treated with BSA, and this increase was attenuated by CerS5 knockdown (Figure 6E). Taken together, all of these results demonstrate a requirement for CerS5 in induction of sphingolipid-mediated autophagic flux by myristate.

Autophagy is required for induction of hypertrophy by myristate in isolated cardiomyocytes. While numerous reports have demonstrated an increase in autophagy in other hypertrophic cardiac conditions, it is still debated whether autophagy promotes or protects against
cardiac hypertrophy in these systems (33–36, 42). To test the role for autophagy in regulation of cardiomyocyte hypertrophy, isolated cardiomyocytes were subjected to LC3B knockdown, which has been shown to impair autophagy (43). Strikingly, knockdown of LC3B prevented myristate-induced cardiomyocyte hypertrophy, suggesting a prohypertrophic role for autophagy in lipid overload (Figure 8). To confirm that this result was not due to an off-target effect of LC3B knockdown, cells were treated with 3-methyladenine, which inhibits autophagy via its action on type III phosphoinositide 3-kinases (44). Consistent with observations with siRNA-mediated knockdown of CerS5. Immunoblot quantitations are presented with representative immunoblots; noncontiguous lanes, separated by white lines, are shown from the same gel. (C) Myristate treatment increased the number of GFP-labeled puncta in GFP-LC3B–expressing cells, suggesting increased numbers of autophagosomes; this effect was attenuated by CerS5 knockdown. All results are presented as mean ± SEM.

Discussion

Despite the considerable work that has been performed to tease apart the mechanisms of lipotoxic DbCM, to date, only general information has been published about the specific lipid mediators governing this process. In particular, although a few studies have implicated ceramides as a lipid class required for induction of cardiotoxicity, the particular species involved have yet to be determined (23, 45–49). Additionally, the practical relevance of lipotoxicity to DbCM remained to be demonstrated, as all effective models of DbCM, to date, relied on genetic manipulations that disrupted cardiac lipid handling in a manner with no clinical parallels (ref 2, reviewed in refs. 3, 4, 22).

The present study addresses both of these concerns. Here, we present a nontransgenic model of diet-induced obesity that robustly produced DbCM-like cardiac hypertrophy and functional impairment. Our findings revealed that an obesogenic diet based on milk fat, rather than lard, induced cardiac dysfunction, both gross and cellular hypertrophy, and increased autophagy in hearts of nontransgenic mice (Figures 1 and 4, and Tables 1 and 2). Furthermore, all of these outcomes were shown to be sphingolipid dependent. This diet, which was highly enriched in myristate (C14:0), also specifically potentiated production of the myristate-containing ceramide species C14:0-ceramide (Figure 2). These findings provided a compelling argument for further in vitro mechanistic work in isolated adult cardiomyocytes, where we were able to identify a specific pathway through which oversupply of an individual saturated fat induced cardiac hypertrophy. In isolated cells, myristate, but not palmitate (C16:0), was found to induce sphingolipid-dependent cardiomyocyte hypertrophy (Figure 3). Furthermore, myristate treatment potentiated C14:0-ceramide production, as was seen in the animal model. Subsequent to these observations, we identified a specific CerS isoform (CerS5) required for induction of hypertrophy by lipid overload (Figure 3).

This work also provided basic insight into the link between autophagy and lipotoxic cardiomyocyte hypertrophy. While numerous reports have demonstrated an increase in autophagy in other hypertrophic cardiac conditions, very few have examined cardiac autophagy in the context of T2D, and none have determined what effect lipid overload has on this pathway. Additionally, some controversy has surrounded the role for autophagy in the hypertrophic heart (31–36, 42). The debate centers on whether autophagy promotes or protects against cardiac hypertrophy in these systems. One major hypothesis unifies these 2 points of view, suggesting that autophagy is a “Goldilocks” phenomenon: too little is pathogenic; too much is pathogenic (50). In an iso-
lated cardiomyocyte system, we demonstrated that oversupply of myristate, but not palmitate, promoted an increase in autophagy (Figures 5 and 6). Furthermore, we directly tested whether inhibition of autophagy would promote hypertrophy or prevent it. In fact, we found that fatty acid–induced hypertrophy was completely prevented by knockdown of LC3B or 3-methyladenine treatment, suggesting a pathogenic role for autophagy in cardiac lipid overload (Figure 8 and Supplemental Figure 8). Finally, we linked the induction of autophagy by myristate to the same CerS isoform implicated in the hypertrophic response (Figures 5 and 6).

Based on these findings, we propose a model in which a diet rich in myristate, which is a major constituent of milk fat–containing products such as butter and ice cream, induces cardiac hypertrophy through its effects on ceramide synthesis and composition. Namely, dietary myristate oversupply promotes ceramide synthesis via CerS5 and, consequently, increased autophagy. This increase in autophagy subsequently promotes hypertrophy through a mechanism that has yet to be identified.

Two major routes could potentially link autophagy and hypertrophy. The first mechanism would be mediated by conventional signaling pathways. While investigations remain ongoing, several potential connections have emerged through bioinformatic analyses. In particular, analysis of protein-protein interaction pathways has revealed links among 3 proteins of the autophagy pathway (i.e., Atg7, Atg12, and RB1CC1/FIP200) and the hypertrophy protein GSK3β via several routes, including FOXO1 and SIRT1 (S.B. Russo, unpublished data). Indeed, literature has already implicated FOXO1 and SIRT1 in regulation of both autophagy and hyper-

Figure 7
Myristate-treated cells do not display decreased autophagosome clearance. After 16 hours of treatment with BSA (A and B) or myristate (C and D), GFP-LC3B–expressing cells were labeled with LysoTracker Red and observed by fluorescence microscopy for 20 minutes. Representative photomicrographs from the beginning and end of the time course (0 minutes and 20 minutes) are shown. Images were taken using a ×60 objective lens. In the panels presented, gamma settings of 0.7 for both red and green channels were used for all images. The heavily stained structure in the upper right quadrants of C and D is an adjacent cardiomyocyte undergoing cell death.
trophy in cardiomyocytes (51–54). Thus, it appears likely that autophagy and hypertrophy pathways may be subject to common regulators and could possibly engage in crosstalk.

The authors also hypothesize a second, less obvious mechanism by which upregulation of autophagy may be required for induction of hypertrophy. To develop the hypertrophic phenotype, cardiomyocytes undergo substantial subcellular remodeling of numerous components, including the mitochondria, sarcoplasmic reticulum, and sarcomeres (55). Autophagy is a key process of cellular remodeling, and in particular, remodeling and recycling of the endoplasmic/sarcoplasmic reticulum and mitochondria (56, 57). On this basis, the authors speculate that, by altering the intracellular landscape via organelle turnover and by recycling intracellular components, induction of autophagy potentiates the organized remodeling prompted by prohypertrophic stimuli.

With regard to the mechanism linking C14-ceramide levels to autophagy, the authors of this study hypothesize that, in addition to upstream signaling roles, the effect of increased C14-ceramide may mediate membrane properties of the autophagosome and, thereby, the activity of membrane-resident proteins involved in autophagy. This notion is based on data from the literature that indicate that sphingolipid N-acyl chain lengths alter the biophysical properties of membranes, including curvature, fluidity, stability, and permeability (58–65). In particular, it has been shown that genetic ablation of CerS2 in mice promoted membrane fusion and budding (66). This was mechanistically attributed to alterations in lipid packing and membrane curvature, which occurred due to a shift toward shorter, unsaturated ceramides and a total ablation of C22- to C24-ceramide production. Furthermore, it has been shown that the activities of membrane-resident proteins are modulated by membrane biophysical properties and are sensitive to membrane lipid composition (67–69). In findings particularly relevant to this study, it was shown that membrane lipid composition and curvature were essential for targeting of Barkor/Atg14 to the budding autophagosome (70). Additionally, it has been shown that the proautophagic activity of BECN1 is mediated by its exit from sphingolipid-enriched membrane microdomains (so-called “lipid rafts”) in the context of traumatic brain injury (71). Thus, it seems plausible that increases in C14-ceramide may induce autophagy by promoting membrane budding and fusion and by altering localization or activity of membrane-resident proteins, including BECN1.

**Methods**

**Diets.** High-fat diets (60% calories from fat) and an isocaloric CD (10% calories from fat; TD.08810) were purchased from Harlan Laboratories Inc. High-fat diets were based on lard (TD.06414) or milk fat (TD.09766). The MFBD used anhydrous milk fat (also known as anhydrous butter oil) as its fat source; this substance is 99.8% fat and contains no detectible carbohydrate, including lactose. Both high-fat diets derived 18.4% of kcal from protein, 21.3% of kcal from carbohydrate, and 60.3% of kcal from fat. All components of the high-fat diets were identical, except for the substitution of anhydrous milk fat for lard. The CD derived 22.3% of kcal from protein, 60.9% of kcal from low-glycemic carbohydrate, and 16.8% of kcal from fat.
**Research Article**

**Animals.** Eight-week-old C57BL/6 male mice (Jackson Laboratory) were given water and chow ad libitum. For the study with time points of 8 and 16 weeks on the diets, 12 mice were placed in each group, and 6 mice were sacrificed per group per time point. For the study with time points of 15 and 18 weeks of feeding, 12 animals were assigned to each group. At the end of the study, at least 7 mice remained per group, with the exception of the lard group that received myricon injections, of which only 5 survived. All mice from this study were sacrificed at 18 weeks. Metabolic characterization of these animals revealed a frank T2D phenotype in the MFBD-fed animals by 8 weeks. In particular, fasting plasma glucose levels in the MFBD mice exceeded 142% of control levels; this is consistent with and exceeds clinical diagnostic guidelines for T2D (73).

**Echo cardiography.** Mice underwent echocardiography to examine in vivo LV structure and function using a 40-MHz transducer and a Vevo 770 echocardiograph (VisualSonics). Three to six beats were averaged for each measurement. LV dimension and wall thickness were made at end-diastole and end-systole using American Society of Echocardiography criteria (74). Mean wall thickness was calculated as the average of interventricular septal wall thickness (IVS) and LV posterior wall thickness (LVPW). Relative wall thickness (RWT) was calculated as the mean LV wall thickness divided by the LV internal dimension at end-diastole (LVIDd). LV mass was calculated as follows: LV mass = 0.8 × 1.05 × [(IVSd + LVPWd + LVIDd)³ – (LVIDd)³]. LV mass was normalized to BW (LV/BW ratio) and tibial length (LV/TL ratio). Ejection fraction (EF, %) was calculated as follows: EF = 100 × (LVEDV – LVESV)/LVEDV, where LVEDV, end-diastolic volume (LVEDV) and end-systolic volume (LVESV) were determined using Simpson’s method of discs. Both systolic (SBP) and diastolic blood pressure (DBP) were measured by using a standard, noninvasive, no-anesthesia Simpson’s method of discs. Both systolic (SBP) and diastolic blood pressure (DBP) were measured by using a standard, noninvasive, no-anesthesia Simpson’s method of discs. Both systolic (SBP) and diastolic blood pressure (DBP) were measured by using a standard, noninvasive, no-anesthesia Simpson’s method of discs. Both systolic (SBP) and diastolic blood pressure (DBP) were measured by using a standard, noninvasive, no-anesthesia Simpson’s method of discs. Both systolic (SBP) and diastolic blood pressure (DBP) were measured by using a standard, noninvasive, no-anesthesia Simpson’s method of discs.

**Cardiomyocytes.** Cardiomyocytes were isolated with TRIzol (Invitrogen) according to the manufacturer’s instructions. Total RNA was isolated with TRIzol (Invitrogen), according to the manufacturer’s protocol. CDNA was synthesized using SuperScript III (Invitrogen). Quantitative real-time PCR was performed using SYBR Green reagent (Invitrogen). Primers for Gapdh, Becn1, Atg7, and Gapdh were obtained from SABiosciences. Feline Cers5 and Becn1 primers were ordered from Integrated DNA Technologies.

**Analysis of cardiomyocyte area.** For analysis of isolated cardiomyocyte areas, cells were treated as indicated, fixed with 10% formalin, stained with H&E, and photographed and analyzed as described above. Mean cell area was determined from 10 fields each from a total of 5 to 6 replicates per group, isolated from a total of 2 separate animals, yielding a final total of 150–200 analyzed cells per treatment group. Cells demonstrating evidence of rounding (an indication of cell death) were excluded from analysis. Data are presented as mean ± SEM. Histograms of cell size, as well as analyses of kurtosis and skewness, are provided in Supplemental Figures 3, 4, and 7. Example photomicrographs showing cell size and morphology are provided in Supplemental Figure 9.

**Western blotting.** Cells were washed twice with sterile filtered cold PBS, lysed with buffer containing 20 mM Tris (pH 7.5), 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 2.5 mM sodium pyrophosphate, 1 mM β-D-glycerophosphate, 1 mM sodium orthovanadate, 1% Triton X-100, and Complete Mini Protease Inhibitor (Roche), disrupted by freeze-thaw, and incubated on ice for 10 minutes. Insoluble material was pelleted by centrifugation at 4°C. Protein concentrations were determined using a Micro BCA Protein Assay Kit (Thermo-Fisher). Lysates were subjected to SDS-PAGE and immunoblotted as described previously (10). Membranes

**Free fatty acid treatment.** To prepare stock solutions of SFAs, medium was supplemented with 2% fatty acid–free BSA and either 1.5 mM myristate or 2.0 mM palmitate. Solutions were sonicated briefly, incubated for 15 minutes at 55°C, and cooled to 37°C. For studies in nontransfected cells, cells were incubated in 0.75 mM myristate or 1.0 mM palmitate for 16 hours. For experiments requiring transfection (described below), cells were treated with 0.1 mM myristate in order to minimize lipotoxicity. For experiments utilizing the de novo sphingolipid synthesis inhibitor myriocin or the autophagy inhibitor 3-methyladenine (Sigma-Aldrich), cells were pretreated with 1.0 μM myriocin, 2.5 mM 3-methyladenine, or vehicle for 30 minutes prior to addition of fatty acids. Inhibitors were allowed to remain in the medium for the duration of fatty acid treatment.

**siRNA transfection and sphingoid base treatment.** Cardiomyocytes were transfected with siRNA oligonucleotides using Lipofectamine 2000 (Invitrogen) according to the manufacturer’s instructions. Custom feline siRNAs directed against CerS5 and LC3B were purchased from Invitrogen. AllStars Negative Control siRNA was from QiAGEN. At 24 hours after transfection, culture medium was supplemented with a final concentration of 2.5 μM C18-DHS (Matreya) or ethanol, or with 0.1 mM myristate or BSA. Cells were treated with DHS for a total of 3 hours or with myristate for a total of 16 hours.

**DNA transfection.** Plasma DNA was transfected into cardiomyocytes using Lipofectamine 2000 (Invitrogen) according to the manufacturer’s protocol. For CerS5 overexpression studies, cells were transfected with empty pcDNA3.0 or pcDNA3.0 containing the human CerS5 cDNA sequence (cloned and validated as described previously, ref. 76) and cultured for 24 hours. The CerS5 construct was provided by Anthony Futerman (Department of Biological Chemistry, Weizmann Institute of Science, Rehovot, Israel). For studies utilizing the GFP-LC3B construct, cells were cotransfected with pSELECT-GFP-LC3 (InvivoGen) and other plasmids or siRNAs, as indicated, and cultured for 24 hours prior to fatty acid treatment, as described above.

**RNA isolation and quantitative real-time PCR analysis.** Total RNA was isolated with TRIzol (Invitrogen), according to the manufacturer’s protocol. CDNA was synthesized using SuperScript III (Invitrogen). Quantitative real-time PCR was performed using SYBR Green reagent (Bio-Rad) as described previously (10). Results were normalized to GAPDH. Primers for Map4k3b, Becn1, Atg7, and Gapdh were obtained from SABiosciences. Feline Cers5 and Becn1 primers were ordered from Integrated DNA Technologies.

**Analysis of cardiomyocyte area.** For analysis of isolated cardiomyocyte areas, cells were treated as indicated, fixed with 10% formalin, stained with H&E, and photographed and analyzed as described above. Mean cell area was determined from 10 fields each from a total of 5 to 6 replicates per group, isolated from a total of 2 separate animals, yielding a final total of 150–200 analyzed cells per treatment group. Cells demonstrating evidence of rounding (an indication of cell death) were excluded from analysis. Data are presented as mean ± SEM. Histograms of cell size, as well as analyses of kurtosis and skewness, are provided in Supplemental Figures 3, 4, and 7. Example photomicrographs showing cell size and morphology are provided in Supplemental Figure 9.

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were incubated with antibodies against BECN1 (Cell Signaling Technology), GFP (Cell Signaling Technology), or actin (Sigma-Aldrich) and with anti-rabbit secondary antibody (Cell Signaling). Proteins were visualized by enhanced chemiluminescence. Images were quantitated using ImageJ software (NIH). Results were presented as mean ± SEM and shown with a representative immunoblot image.

Fluorescence microscopy of cardiomyocytes. After transfection and fatty acid treatment (described above), half of the culture medium was removed and saved, and cells were preloaded for 30 minutes with 50 nM Lysotracker Red DND-99 (Invitrogen). The dye-containing medium was then removed and replaced with the dye-free aliquot, and cells were examined by confocal fluorescence microscopy using an Olympus FV10i microscope. To track autophagosome dynamics, images were taken every 5 minutes for 20 minutes at a magnification of ×60 with a zoom setting of x39. Identical laser and sensitivity settings were utilized for all images. Images were analyzed using FV10-ASW software, as described above, using identical threshold and gamma settings for images presented in figures. For quantification of GFP-labeled puncta, cells were formalin fixed in the dark, rinsed in PBS, photographed at a magnification of ×60 with a zoom setting of ×2 and analyzed with FV10-ASW software, as described above. Puncta were counted for 25 cells per treatment group. Results are presented as mean ± SEM.

Statistical analysis. Data are expressed as mean ± SEM. Animal data were analyzed by 1-way ANOVA followed by Bonferroni’s test. All other data were analyzed by 2-tailed, unpaired Student’s t-test. For all experiments, P < 0.05 was considered statistically significant.

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