Glucose and palmitate uncouple AMPK from autophagy in human aortic endothelial cells

Karen A. Weikel, José M. Cacicedo, Neil B. Ruderman, and Yasuo Ido

Department of Medicine, Boston University School of Medicine and Boston Medical Center, Boston, Massachusetts

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Weikel KA, Cacicedo JM, Ruderman NB, Ido Y. Glucose and palmitate uncouple AMPK from autophagy in human aortic endothelial cells. Am J Physiol Cell Physiol 308: C249–C263, 2015. First published October 29, 2014; doi:10.1152/ajpcell.00265.2014.—Dysregulated autophagy and decreased AMP-activated protein kinase (AMPK) activity are each associated with atherogenesis. Atherogenesis is preceded by high circulating concentrations of glucose and fatty acids, yet the mechanism by which these nutrients regulate autophagy in human aortic endothelial cells (HAECs) is not known. Furthermore, whereas AMPK is recognized as an activator of autophagy in cells with few nutrients, its effects on autophagy in nutrient-rich HAECs has not been investigated. We maintained and passaged primary HAECs in media containing 25 mM glucose and incubated them subsequently with 0.4 mM palmitate. These conditions impaired basal autophagy and rendered HAECs more susceptible to apoptosis and adhesion of monocytes, outcomes attenuated by the autophagy activator rapamycin. Glucose and palmitate diminished AMPK activity and phosphorylation of the uncoordinated-51-like kinase 1 (ULK1) at Ser555, an autophagy-activating site targeted by AMPK. 5-Aminimidazole-4-carboxamide-1-β-d-ribofuranoside (AICAR)-mediated activation of AMPK phosphorylated acetyl-CoA carboxylase, but treatment with AICAR or other AMPK activators (A769662, phenformin) did not restore ULK1 phosphorylation or autophagosome formation. To determine whether palmitate-induced ceramide accumulation contributed to this finding, we overexpressed a ceramide-metabolizing enzyme, acid ceramidase. The increase in acid ceramidase expression ameliorated the effects of excess nutrients on ULK1 phosphorylation, without altering the effects of the AMPK activators. Thus, unlike low nutrient conditions, AMPK becomes uncoupled from autophagy in HAECs in a nutrient-rich environment, such as that found in patients with increased cardiovascular risk. These findings suggest that combinations of AMPK-independent and AMPK-dependent therapies may be more effective alternatives than either therapy alone for treating nutrient-induced cellular dysfunction.

endothelium; autophagy; palmitate; glucose; AMPK

Macrouathagy (hereinafter referred to as “autophagy”) is a process by which cellular components are engulfed in a double-membraned vesicle, the autophagosome, and subsequently degraded by lysosomal proteases (17). Autophagy plays a key role in the cellular stress response, in which it can be induced by conditions such as starvation, inflammation, and oxidative stress (19, 51, 78). Regulation of starvation-induced autophagy is well-studied and involves complex signaling pathways that are modulated by the activities of several kinases including mammalian target of rapamycin complex 1 (mTORC1), uncoordinated-51-like kinase 1 (ULK1), and AMP-activated protein kinase (AMPK) (69, 89). Under low nutrient conditions, these kinases (among others) coordinate the formation of an autophagosome, to which autophagic substrates are recruited by adaptor proteins such as p62 (Fig. 1) (45). The autophagosome then fuses with a lysosome to create an autolysosome in which proteases, such as cathepsins, digest the target substrates along with their adaptor proteins (Fig. 1). Their breakdown products are then used to synthesize new proteins or fuel the cell (43).

A much less understood aspect of autophagy is regulation of its basal activity. Basal autophagy rids the cell of aged and damaged organelles when nutritional supply is not limited and is critical for cellular homeostasis in the vasculature as has been demonstrated by ApoE-null mice fed a Western diet (a model for atherosclerosis). Compared with wild-type mice, basal autophagy is impaired in the aorta of ApoE-null mice and its impairment in macrophages is associated with increased plaque development (74). These findings suggest that basal autophagy may be altered during atherosclerosis, but it remains to be determined how this type of autophagy is regulated and whether the same regulatory kinases and stimuli that modulate starvation-induced autophagy are involved. Importantly, it is not clear how hyperglycemia and dyslipidemia (specifically elevated free fatty acids), as observed in patients with type 2 diabetes and the metabolic syndrome, modulate basal autophagy in endothelial cells, the site of atherogenesis (3). Previous work in the pancreatic β-cell and macrophage has linked high concentrations of glucose and palmitate [a saturated fatty acid that has been associated with cellular toxicity (8, 9)] with autophagy inhibition (58, 88), but the ways in which these nutrients regulate basal autophagy in vascular tissues, such as human aortic endothelial cells (HAECs), have not been well characterized.

AMPK is a serine/threonine kinase that stimulates catabolic and inhibits anabolic processes to restore ATP levels when cellular energy is low (66). Activation of AMPK attenuates endothelial cell dysfunction (20) and aortic lesion formation (86) and also promotes starvation-induced autophagy through interactions with mTORC1 and/or ULK1 (Fig. 1) (21, 29, 42). It is not known, however, whether AMPK regulates basal...

Address for reprint requests and other correspondence: K. A. Weikel, Dept. of Medicine, Boston Univ. School of Medicine and Boston Medical Center, 650 Albany St., Rm. 820, Boston, MA, 02118 (e-mail: karen.weikel@bmc.org).

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C249
autophagy in the excess nutrient environment that predisposes endothelial cells to dysfunction and atherogenesis. Elucidation of this nutrient-AMPK-autophagy relationship in HAECS could be especially important since decreases in both AMPK activity and autophagy have been associated with an increased risk of atherosclerosis (20, 74, 86). An understanding of how hyperglycemia and dyslipidemia affect endothelial cell basal autophagy as well as the role of AMPK in this regulation could be critical for the development of pharmacological therapies that could target the endothelium to prevent atherogenesis. Thus, in the present study we first characterized autophagy in cultured primary HAECS incubated in high concentrations of glucose and palmitate, and then evaluated its regulation by AMPK.

**EXPERIMENTAL PROCEDURES**

**Materials.** Human aortic endothelial cells (HAECS), human umbilical vein endothelial cells (HUVECs), and EGM-2 media (catalog no. cc-3162) were purchased from Lonza (Walkersville, MD). HEPES-buffered phenol red free media 199 (catalog no. M-2520, Sigma, St. Louis, MO), was supplemented with 5% FBS (catalog no. 26140-087, Life Technologies, Carlsbad, CA) and 50 μM carmine (catalog no. C-0283, Sigma). Bafilomycin (catalog no. B1793), phenformin hydrochloride (catalog no. P7045), oleate (catalog no. O1008), and chloroform (catalog no. C7559) were also purchased from Sigma. Palmitate was purchased from Nu-Chek Prep (Elysian, MN). 5-Aminomimidazole-4-carboxamide-1-β-D-ribofuranside (AICAR; catalog no. A611700) was purchased from Toronto Research Chemicals (North York, Ontario, Canada), and rapamycin (catalog no. R-5000) was purchased from LC Laboratories (Woburn, MA). A769662 (catalog no. 3336) was purchased from Tocris Biosciences (Minneapolis, MN). p62 antibody (catalog no. 610832) was purchased from BD Biosciences (San Jose, CA). Antibodies for LC3-II (catalog no. 38685), phosphorylated (Thr172) AMPKα1/2 (catalog no. 2531S), phosphorylated (Ser79) acetyl-CoA carboxylase (ACC; catalog no. 3661S), ACC (catalog no. 4190S), phosphorylated (Thr 389) p70S6K (catalog no. 9205S), phosphorylated (Ser2448) mTOR (catalog no. 2971S), and phosphorylated (Ser555) ULK1 (catalog no. 5869S) were purchased from Cell Signaling Technology (Danvers, MA). Total AMPKα1 (catalog no. 3694-1) was purchased from Epionce (Burlingame, CA). Total ULK1 (catalog no. sc-33182), mTOR (catalog no. 2972S), GAPDH (catalog no. sc-25778), and acid ceramidase (catalog no. sc-28486) antibodies were purchased from Santa Cruz Biotechnology (Dallas, TX). Actin antibody (catalog no. A4700) was purchased from Sigma (St. Louis, MO). Methanol (catalog no. A452-4) and acetic acid (catalog no. A38-212) were purchased from Fisher Scientific (Pittsburgh, PA). Secondary peroxidase-conjugated anti-mouse (catalog no. NAX931) and anti-rabbit (catalog no. NA934V) antibodies were purchased from GE Healthcare (Pittsburgh, PA). 3P and [3H]-L-serine were purchased from Perkin Elmer (Boston, MA).

**Cell culture.** HAECS were grown in EGM-2 media containing 2% FBS on Primaria culture dishes (reference no. 353803; BD Biosciences). For hyperglycemic conditions, cells were grown and passaged in EGM-2 media supplemented with 1-glucose (catalog no. G-7021) (Sigma) to a final concentration of 25 mM. These cells were incubated in hyperglycemic media beginning in passage 3 and when they reached 85% confluence were split in hyperglycemic media for passage 4. This protocol of maintaining and splitting the cells in hyperglycemic media was repeated for future passages. Because we are interested in modeling the long-term effects of hyperglycemia, rather than the transition to hyperglycemia, experiments using hyperglycemic cells began once they reached passage 4. At this stage, all of the cells were generated in a hyperglycemic environment and therefore exposed to it for their entire lifetime.

For both normoglycemic and hyperglycemic cells, experiments were performed when cells were 85–95% confluent. When applicable, HAECS were pretreated with the indicated concentrations of AICAR, A769662, phenformin, or rapamycin in EGM-2 for 30 min. Medium was then changed to DMEM (catalog no. 31600-034, Life Technologies) supplemented with 5% FBS containing either BSA, BSA-palmitate or BSA-oleate conjugates [3:1 free fatty acid (FFA):BSA ratio], as described previously (9). Briefly, BSA was used to both stabilize the insoluble fatty acids and transport them to the cell. Control-treated cells were exposed to media containing 0.5 mM BSA, while fatty acid-treated cells were exposed to media containing BSA conjugated to 0.4 mM palmitate or oleate at a 3:1 FFA:BSA molar ratio. Cells were exposed to BSA or BSA-fatty acid conjugates in DMEM for 6 h prior to harvesting for Western blotting, mRNA analysis, cathepsin L activity, and monocyte adhesion. For those experiments that included AICAR, A769662, phenformin, or rapamycin, these compounds were also present in the DMEM for the 6 h incubation. Prior to TUNEL staining, cells were exposed to experimental conditions in DMEM for 16 h.

**Western blotting.** Cells were harvested and lysed in a Triton-based buffer containing 20 mM Tris pH 7.5, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 0.1% SDS, 1% Triton X-100, 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, 1 mM sodium orthovanadate, and 1 μg/ml leupeptin. Proteins were separated using either 4–12% Bis-Tris gradient or 14% Tris-glycine gels (Life Technologies) and transferred to PVDF membranes. After blocking for at least 1 h at room temperature with nonfat milk, membranes were incubated in primary antibody overnight. Following washing in TBST, membranes were incubated in peroxidase-conjugated anti-mouse, anti-rabbit, or anti-goat antibodies and bands were visualized on film using SuperSignal West Pico (catalog nos. 1856135 and 1856136) or Femto (catalog no. 34095) chemiluminescent substrate (Thermo Scientific, Rockford, IL). Densitometric analyses were carried out using Scion Image software and are presented after adjustment for expression of loading controls (actin or GAPDH).

**Assessment of autophagosome formation.** Cells were exposed to experimental conditions as described above for 6 h, in the presence or absence of 10 mM bafilomycin. Bafilomycin-induced LC3-II protein levels, or the ratio of LC3-II/LC3-I protein, as measured with Western blotting, were used as indicators of autophagosome abundance. To measure LC3-II protein accumulation by fluorescence microscopy, HAECS were infected with green fluorescent protein (GFP)-LC3-expressing lentivirus. Before comparing LC3-II puncta between treatments, we first set a threshold of “autophagy-induced LC3 accumulation,” defined as the average pixel fluorescence in a single cell treated with rapamycin. Puncta fluorescence was visualized on a Nikon TE-200 Eclipse Inverted microscope (Melville, NY) and quantified with ImageJ (National Institutes of Health, Bethesda, MD). We then calculated the percentage of cells in each treatment group (control, control+baflomyacin, nutrient excess, nutrient excess+baflomyacin, nutrient excess+rapamycin, nutrient excess+rapamycin+baflomyacin, nutrient excess+AMPAK activator, nutrient excess+AMPAK activator+baflomyacin) that reached or surpassed this threshold. Bafilomycin-induced change in the percentage of cells above threshold was calculated for nutrient excess, nutrient excess+rapamycin and nutrient excess+AMPAK activator conditions relative to control treatment.

**Recombinant adenosine and lentivirus.** The adenovirus expressing a constitutively active mutant of AMPK (α1/1, T172D) was constructed from a truncated rat α1-subunit with aspartate substituted for threonine 172, as described previously (41, 90). The adenosine-expressing ceramidase was created as previously described (8). Lentivirus-expressing acid ceramidase (ASAHI, accession no. BC016481) was created by LR reaction of pENT221 (HisCD00039940, Harvard Plasmid) with a home-made lentivirus vector (containing a gateway cassette) to produce...
a His-Flag tag at the COOH terminus of the protein. Infection of HAECs with adenovirus-expressing acid ceramidase had similar effects as infection with lentivirus-expressing acid ceramidase. The recombinant lentivirus plasmids expressing β-galactosidase and GFP-LC3 (Addgene no. 22418) were created in a home-made Gateway compatible lentivector after subcloning them into pENTR1A vector. All lentiviruses were produced as described previously (55).

**Cathepsin L activity.** Cathepsin L activity was measured using the Magic Red Cathepsin L Assay kit (catalog no. 941) purchased from Immunochrometry Technologies (Bloomington, MN) by following the manufacturer’s instructions. Briefly, HAECs were treated with the indicated concentrations of glucose and palmitate along with a cresyl violet fluorophore that becomes fluorescent upon activation of cathepsin L. Cathepsin L activity was quantified on a Tecan microplate reader (San Jose, CA) and relative change in activity was calculated after adjusting for cell density and the number of acidic organelles.

**AMPK activity assay.** AMPK activity was measured as reported previously (30). In this assay, γ-32P (catalog no. NEX011001MC, Perkin Elmer), ATP and SAMs peptide (HMRSMASGLHVLKKR) were incubated with cell lysate. AMPK activity was assessed by measuring radioactive counts of 32P following the reaction.

**TUNEL staining.** Cells were labeled using Roche’s In Situ Cell Death Detection Kit, AP (catalog no. 11 684 809 910) (Indianapolis, IN) according to the manufacturer’s instructions. TUNEL-positive staining was visualized using NBT-BCIP substrate (catalog no. 11 681 451 001) (Roche, Indianapolis, IN), and pictures were obtained on a Nikon TE-200 Eclipse Inverted microscope.

**Real-time PCR.** SYBR green master Mix from Clontech (catalog no. RR420A; Mountain View, CA) was used to quantify real-time PCR products on Cepheid's Smart Cycler (Hamburg, PA). The following primers were used to detect mRNAs: β-actin: 5′-TTGTAAACCA-ACTGGGAGCATATGG-3′ (sense) and 5′-GATCTTGATCTTCT-CAT-GGTGCTAGG-3′ (anti-sense); E-selectin: 5′-CTCTGACAGAAG-AGCCAAG-3′ (sense), 5′-ACTTTGATCTCAGTGAAGCCA-3′ (anti-sense). mRNA levels of E-selectin were quantified using the 2−ΔΔCT method, relative to actin.

**Monocyte adhesion.** THP-1 monocytes (human cell line) were grown in RPMI-1640 media (catalog no. R-1383; Sigma) supplemented with 10% FBS. Monocytes were labeled with carboxyfluorescein succinimidyl ester (CFSE) dye (catalog no. C1157) (Life Technologies) and incubated with unsupplemented RPMI-1640 media for 30 min. Labeled monocytes were then incubated with HAECs for 1 h at 37°C. Unattached monocytes were washed off of HAECs with PBS, and the number of monocytes adherent to the HAECs were quantified using a microplate reader (Tecan).

**Measurement of ceramide and sphingosine.** HUVECs were incubated in M199 media containing either 5 or 30 mM glucose and either 0 or 0.3 mM palmitate (as indicated in Fig. 6a) as well as 2 mM/ml [3H]-l-serine for 16 h at 37°C. After measurement of specific activity in 100 μl of media, cells were washed three times with cold PBS. PBS (0.9 ml) containing 0.2% SDS was then added to each well, 100 μl of which was used to determine protein concentration. The remainder was transferred to fresh tubes, to which 3 ml CHCl3:MeOH (1:2, vol/vol) was added. One milliliter each of chloroform and water was then added and the sample was vortexed for 1 min. Samples were centrifuged for 5 min at 2,000 rpm to separate the phases, and the aqueous layer was aspirated while the organic phase was transferred to a new tube. Lipids were dried and resuspended in 50 μl CHCl3:MeOH (1:1). Ten micrograms of each standard were added to each tube and loaded onto a thin-layer chromatography plate and run in the solvent system [CHCl3:MeOH:3.5% N NH4OH (85:15:1)]. The bands were exposed to iodine and scraped into scintillation vials containing 1 ml MeOH:acetic acid (80:20). Samples were sonicated until bands dissolved. Four milliliters sonication fluid were added, and [3H] was measured on a scintillation counter (3 min/vial).

**Statistics.** Data are presented as means ± SE. Unless stated otherwise, differences between treatment groups were analyzed using ANOVA, adjusting for multiple comparisons (GraphPad software, La Jolla, CA). P < 0.05 was considered significant.

**RESULTS.** In patients with poorly controlled type 2 diabetes, blood glucose levels are consistently elevated while fatty acid levels rise to peak concentrations during the 6–8 h of overnight fasting (27, 32, 68, 73, 75). Compared with control patients, free fatty acid levels in such patients with diabetes are 50% higher during overnight fast (68). Postprandial levels of palmitate (one of the predominant saturated fatty acids) in these patients have been reported to be nearly three times higher than healthy, nondiabetic subjects (68). To mimic this constant exposure of aortic endothelial cells to glucose interspersed with a subacute fatty acid (palmitate) exposure, we utilized a unique cell culture model to better understand the pathological effects of both of these nutrients on HAECs.

In our model, we sought to capture the long-term cellular effects of hyperglycemia, rather than the effects of a transition from normoglycemic (5 mM glucose) to hyperglycemic conditions (25 mM glucose). Therefore, HAECs destined for treatment with 25 mM glucose were switched from normoglycemic to hyperglycemic media beginning in passage 3. This medium was used for growth of these cells as well as their subsequent splitting into future generations. Experimental treatment and analysis of these cells began once the HAECs reached 95% confluence in passage 4 (~10 days after beginning the use of the hyperglycemic media) and continued in future passages. Similar to the aortic endothelial cells of patients with poorly controlled type 2 diabetes, all of the cells beginning in the fourth passage of our model were generated in a hyperglycemic environment and therefore exposed to this stress for their entire lifetime (“chronic exposure”).

To model the acute surges in palmitate experienced by patients with diabetes, we treated HAECs grown under hyperglycemic conditions with 0.4 mM palmitate [a physiologically relevant concentration that can be observed in diabetes patients during fasting or exercise (32, 75, 76)] for 6 h (“acute” exposure). Significant effects of palmitate were not observed with a shorter treatment (3 h), and longer incubations (24 and 48 h) did not augment the effects observed after 6 h.

**Glucose with palmitate decreases autophagic flux in primary human aortic endothelial cells.** Incubation of endothelial cells with high concentrations of glucose and fatty acids (including palmitate) has been shown to increase oxidative stress and inflammation (9, 41, 64), but their effects on autophagy in HAECs are not known. We began our investigation in HAECs by measuring protein levels of p62, a substrate adaptor protein that is degraded with its cargo by the autolysosome (Fig. 1). Thus, p62 protein accumulates when autophagy is impaired.

Compared with HAECs incubated in control conditions (5 mM glucose, 0 mM palmitate), cells incubated in media with excess nutrients (chronic exposure to hyperglycemic media followed by a 6-h treatment with 0.4 mM palmitate) accumulated significantly more p62 protein (Fig. 2). Preliminary data showed that either chronic exposure to glucose or acute exposure to palmitate also increased p62 protein levels, although not as robustly as when the nutrients were combined. Therefore, subsequent analyses of autophagy presented here were per-
formed under the stress of both nutrients. Pretreatment with rapamycin, an inhibitor of mTORC1 (an enzyme that inhibits autophagy), ablated the nutrient-induced accumulation of p62, suggesting that under nutrient excess conditions, autophagy is impaired (Fig. 2A). Along with the saturated fatty acid palmitate, oleate, a monounsaturated fatty acid, is one of the most abundant fatty acids in circulation. We observed that HAECs exposed to elevated concentrations of glucose and oleate, instead of palmitate, did not accumulate p62, suggesting that all fatty acids do not impair autophagy (data not shown).

Glucose with palmitate impairs autophagosome formation and cathepsin L activity. Autophagosome formation, a fundamental step in autophagy (Fig. 1, panels 1 and 2), can be quantified indirectly by measuring the protein levels of microtubule-associated protein light chain 3 (LC3), a membrane-bound component of the autophagosomal membrane (Fig. 1) (89). LC3 is present in the cytosol as LC3-I (16 kDa), but upon conjugation with phosphatidylethanolamine it is incorporated into the developing autophagosome membrane as LC3-II (Although the molecular weight of LC3-II exceeds that of LC3-I, LC3-II appears on SDS-PAGE at ~14 kDa due to its increased hydrophobicity.). Protein levels of LC3-II, or the LC3-II/LC3-I ratio (indicative of incorporation into the developing autophagosome) are surrogate indicators of autophagosome abundance and formation. Since LC3-II is continually turned over during autophagy, these measurements need to be performed in the presence of an autophagic inhibitor such as bafilomycin A1 (hereinafter referred to as simply bafilomycin) that prevents LC3-II degradation (50, 52). Thus, treatment with bafilomycin causes an accumulation of LC3-II protein. A decrease in bafilomycin-induced LC3-II protein accumulation would suggest that autophagosome formation is diminished.

We infected HAECs with lentivirus-expressing GFP-LC3 and measured LC3 puncta (indicative of autophagosomes) (50) in the presence of bafilomycin, using fluorescence microscopy. LC3 puncta were abundant in HAECs incubated in control conditions, but very few puncta accumulated in HAECs incubated in the presence of excess nutrients (Fig. 2B). Pretreatment of such HAECs with rapamycin restored LC3 puncta to levels observed in the control-treated cells (Fig. 2B). To confirm these findings, we used Western blotting to measure bafilomycin-induced LC3-II protein levels and the LC3-II/LC3-I ratio in HAECs incubated in control and nutrient excess conditions (Fig. 2C). Compared with control-treated cells, cells exposed to a nutrient excess showed decreased protein levels of both LC3-II and the LC3-II/LC3-I ratio (Fig. 2C, lane 4 vs. lane 2). Similar to the data observed by microscopy, rapamycin treatment prevented nutrient-induced decreases in LC3 protein (Fig. 2C, lane 6 vs. lane 4).

Following autophagosome formation, the autophagosome fuses with the lysosome to create an autolysosome in which p62-substrate complexes and LC3-II are degraded by proteases (depicted as scissors in box) (89).

Fig. 1. Simplified model of starvation-induced autophagy. Panel 0: when mammalian target of rapamycin complex 1 (mTORC1) is active, it maintains uncoordinated-51-like kinase 1 (ULK1) in a hyperphosphorylated state that suppresses ULK1 activity and autophagy (89). Panel 1: activation of AMP-activated protein kinase (AMPK) under low nutrient conditions has been associated with phosphorylation of the raptor component of mTORC1, facilitating dissociation of mTORC1 from the ULK1 complex and activation of ULK1 (29, 42). AMPK can also activate ULK1 by phosphorylating Ser555 (21). Panel 2: ULK1 then translocates to the site of autophagosome membrane biogenesis. As the autophagosomal membrane elongates, microtubule-associated protein light chain 3 (LC3)-II binds to the inner membrane of the forming autophagosome and tethers p62 in complex with its substrate. Panel 3: once the autophagosome is formed, it fuses with the lysosome to create an autolysosome in which p62-substrate complexes and LC3-II are degraded by proteases (depicted as scissors in box) (89).
As shown in Fig. 3B, also assessed the adhesion of THP-1 monocytes to HAECs. Changes in E-selectin transcription might recruit more mononuclear cells to the endothelium, thus predisposing changes in E-selectin, a proinflammatory cytokine that facilitates the recruitment of mononuclear cells to the endothelium (23). Expression of E-selectin mRNA in cells incubated in a hyperglycemic media containing palmitate was impaired in HAECs incubated in nutrient excess conditions, because mTORC1 activity is increased. However, we did not

Excess nutrients uncouple AMPK from autophagy

**C253**

Glucose and palmitate decrease AMPK activity in HAECs.

The data presented in Figs. 1–3 demonstrate that incubation of HAECs in high concentrations of palmitate and glucose decreases autophagy, increases inflammation, and predisposes the cells to apoptosis. Furthermore, both the inflammation and apoptosis were attenuated by rapamycin, an mTORC1 inhibitor that induces autophagy. This suggested that autophagy may be impaired in HAECs incubated in nutrient excess conditions because mTORC1 activity is increased. However, we did not

thelial dysfunction, inflammation, in HAECs exposed to excess nutrients. For this purpose, we measured the expression of E-selectin, a proinflammatory cytokine that facilitates the recruitment of mononuclear cells to the endothelium (23). Expression of E-selectin mRNA in cells incubated in a hyperglycemic media containing palmitate was slightly higher (albeit nonsignificantly, \( P = 0.07 \)) than in control-treated cells, a response that was ablated by treatment with rapamycin (Fig. 3B). To determine whether these changes in E-selectin transcription might recruit more mononuclear cells to the endothelium, thus predisposing HAECs to inflammation and potential atherogenesis, we also assessed the adhesion of THP-1 monocytes to HAECs. As shown in Fig. 3B, incubation of HAECs in media containing excess glucose and palmitate increased the number of monocytes adherent to the HAECs, a clear indication that the endothelial cells became inflamed. Similar to its effects on E-selectin, rapamycin prevented this adhesion (Fig. 3B).

Glucose and palmitate decrease AMPK activity in HAECs. The data presented in Figs. 1–3 demonstrate that incubation of HAECs in high concentrations of palmitate and glucose decreases autophagy, increases inflammation, and predisposes the cells to apoptosis. Furthermore, both the inflammation and apoptosis were attenuated by rapamycin, an mTORC1 inhibitor that induces autophagy. This suggested that autophagy may be impaired in HAECs incubated in nutrient excess conditions because mTORC1 activity is increased. However, we did not

Fig. 2. Exposure to high concentrations of glucose and palmitate impairs autophagy. A: human aortic endothelial cells (HAECs) incubated in media containing high concentrations of glucose and palmitate increased levels of p62 protein, an autophagic substrate. Rapamycin, a chemical inducer of autophagy, attenuated p62 protein levels. A representative Western blot of lysates is shown for p62 and actin below a densitometric comparison of p62 protein levels (adjusted for cell density and the number of acidic organelles (\( n = 3 \)). The data presented in Figs. 1–3 demonstrate that incubation of HAECs in high concentrations of palmitate and glucose decreases autophagy, increases inflammation, and predisposes the cells to apoptosis. Furthermore, both the inflammation and apoptosis were attenuated by rapamycin, an mTORC1 inhibitor that induces autophagy. This suggested that autophagy may be impaired in HAECs incubated in nutrient excess conditions because mTORC1 activity is increased. However, we did not

![Fig. 2. Exposure to high concentrations of glucose and palmitate impairs autophagy.](image)

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observe any increase in protein levels of phosphorylated (Thr389) p70S6K nor the ratio of phosphorylated (Ser2448) mTOR/total mTOR protein (data not shown), two downstream targets of active mTORC1. Thus, we focused our attention on another kinase that has been reported to induce autophagy (at least under low nutrient conditions), AMPK. We hypothesized that autophagy impairment observed in the presence of high concentrations of glucose and palmitate also increased mRNA levels of the proinflammatory cytokine E-selectin (n = 3) and adhesion of THP-1 monocytes (n = 9) relative to control-treated cells. Treatment with rapamycin attenuated these changes. For A and B, +P = 0.07 and *P ≤ 0.05 compared with control conditions; **P < 0.05 compared with excess nutrient conditions.

Activation of AMPK does not restore autophagy in a nutrient-laden environment. To address whether AMPK activity regulates nutrient-induced changes in autophagy in HAECs, we treated HAECs with AICAR, a chemical activator of AMPK, under control and excess nutrient conditions (81). We found that under both of these conditions, AICAR activated AMPK. AICAR increased protein levels of phosphorylated AMPK (Thr172) and also increased protein levels of phosphorylated acetyl-CoA carboxylase (ACC) at Ser79, a downstream target of AMPK (49) (Fig. 4, C and D). Since AICAR restored AMPK phosphorylation as well as that of its downstream target under nutrient excess conditions, we measured LC3 protein levels to determine whether autophagosome formation was also restored. Surprisingly, in the presence of high concentrations of glucose and palmitate, AICAR did not attenuate the loss of bafilomycin-induced LC3-II protein (Fig. 5, A and B, lane 8 vs. lane 6). Similarly, it did not alleviate the effects of glucose and palmitate on the ratio of LC3-II/LC3-I or the appearance of GFP-LC3 puncta (data not shown). Unlike AICAR, which is metabolized to an AMP mimic that activates AMPK, A769662 is an allosteric activator of AMPK (16). Treatment of HAECs with 10 μM A769662 increased phosphorylation of AMPK at Thr172 (data not shown), but like AICAR it did not restore LC3-II protein (Fig. 5, A and C, lane 8 vs. lane 6), the ratio of LC3-II/LC3-I (data not shown), nor GFP-LC3 puncta (data not shown) in excess nutrient-exposed cells. Another chemical activator of AMPK, phenformin, is a biguanide and an analog of the diabetes drug metformin (33). As expected, treatment of HAECs with 1 mM...
phenformin increased phosphorylation of AMPK at Thr172 (data not shown). Similar to the effects observed with AICAR and A769662, phenformin treatment did not significantly attenuate nutrient-induced changes in LC3 protein levels (Fig. 5, A and D, lane 8 vs. lane 6 and data not shown). Nutrient-induced changes in LC3 protein levels were also not affected by infection of HAECs with an adenovirus-expressing (α1/α2) ULK1 can be inactivated via mTORC1-dependent phosphorylation (Thr172) (41, 90). Compared with activation of AMPK by AICAR, a mutant form of AMPK that is constitutively active (data not shown) (41, 90). Compared with activation of AMPK by chemical activators, AMPK activation with this mutant was minimal.

mTORC1, AMPK, and ULK1 interact in a complex, dynamic relationship to modulate starvation-induced autophagy (1). ULK1 can be inactivated via mTORC1-dependent phosphorylation (Fig. 1, panel 0). When mTORC1 activity declines, it dissociates from ULK1, perhaps enabling autophagy-inducing kinases such as AMPK to activate ULK1 (Fig. 1, panel 1). A number of studies have shown that Ser555 on ULK1 is a target of AMPK and that its phosphorylation is required for starvation-induced autophagy (21, 29, 48, 60, 88). To address the dissociation between AMPK activation and autophagosome formation in HAECs exposed to nutrient excess conditions, we measured the effect of these nutrients on phosphorylation of Ser555 on ULK1. We found that, compared with control-treated cells, HAECs exposed to nutrient excess conditions had lower protein levels of phosphorylated ULK1 (Ser555) (Fig. 5E). This suggests that in HAECs, phosphorylation of ULK1 at Ser555 is an important feature of basal autophagy induction. In addition to Ser555, we found that incubation of cells with glucose and palmitate also decreased ULK1 phosphorylation at Ser317, another target of AMPK (data not shown) (48). Treatment of nutrient-laden HAECs with AICAR, A769662, or phenformin to activate AMPK did not restore ULK1 phosphorylation at Ser317 (data not shown) (48). Treatments of nutrient-laden HAECs with AICAR, A769662, or phenformin to activate AMPK did not restore ULK1 phosphorylation at Ser555 (Fig. 5E) nor did expression of a constitutively active AMPK (data not shown) (48). A representative Western blot is shown below the graph. D: densitometric analysis of the ratio of phosphorylated (Ser79) acetyl-CoA carboxylase (p-ACC)/total ACC (t-ACC) protein (a target of active AMPK) indicates that aminoimidazole-4-carboxamide-1-β-4-ribofuranoside (AICAR) increases AMPK phosphorylation (n = 5). A representative Western blot is shown below the graph. For A and B, *P < 0.05 compared with control treatment, and for C and D, *P < 0.05 for the effect of AICAR (two-way ANOVA). Overexpression of acid ceramidase restores nutrient-induced losses in ULK1 phosphorylation at Ser555. To address the question of why AMPK is uncoupled from autophagy signaling in the presence of excess glucose and palmitate, we drew on our observation that palmitate, but not oleate, exposure was associated with impaired autophagy (as indicated by an accumulation of p62 protein) (Fig. 2A and data not shown). One of the differences between these two fatty acids is that only palmitate contributes to the cellular synthesis of ceramide, a lipid species that has been shown to decrease AMPK activity and contribute to vascular dysfunction (91, 94). Ceramide and its metabolite, sphingosine-1-phosphate, have been suggested to have opposing roles in autophagy regulation, but whether ceramide accumulation induces or impairs basal autophagy in HAECs remains unclear (70, 84). Since ceramide may potentially regulate both AMPK and autophagy, we hypothesized that, in the presence of palmitate, ceramides may prevent...
activated AMPK from phosphorylating ULK1 and inducing autophagy.

Previous work from our laboratory indicated that palmitate treatment increased ceramides in retinal pericytes (8). As an extension of this finding, we sought to determine whether macrovascular endothelial cells had a similar response. To perform the radioactive tracer experiments and thin-layer chromatography required to resolve ceramides and their related lipid intermediates, a very large number of cells was required per sample. This posed a challenge since HAECs are notoriously poor growers. Thus, we used HUVECs, which can be readily expanded to large quantities. We found that incubation of HUVECs in 0.3 mM palmitate increased ceramide generation (Fig. 6A). Furthermore, ceramide generation also increased in HUVECs incubated in 30 mM glucose or a combination of glucose with palmitate (Fig. 6A). Ceramides are utilized by many cells for the synthesis of a number of different structural and signaling lipid species including sphingosine (the precursor to sphingosine-1-phosphate), glucosylceramides, and sphingomyelin (57). We also found this to be the case in HUVECs, as high concentrations of glucose and palmitate increased intracellular generation of sphingosine (Fig. 6A). Conversion of ceramide to sphingosine is catalyzed by a lysosomal enzyme, acid ceramidase (57). Overexpression of acid ceramidase prevents palmitate-induced accumulation of ceramides (14), while its impairment results in ceramide accumulation (38). To assess the potential role of nutrient-induced ceramides in modulation of the phosphorylation of ULK1, we ectopically expressed acid ceramidase in HAECs (Fig. 6B).
Unlike noninfected (and β-galactosidase-infected) HAECs in which glucose and palmitate treatment decreased phosphorylation of ULK1 at Ser555 (Fig. 5E and data not shown), HAECs overexpressing acid ceramidase had higher levels of phosphorylated ULK1 when incubated with excess glucose and palmitate (Fig. 6C and data not shown). In this nutrient-laden context, treatment of these cells with AICAR or A769662 did not further increase expression of phosphorylated ULK1 (Fig. 6D). These data suggest that, in the presence of glucose and palmitate, accumulation of ceramides (which would be diminished by acid ceramidase overexpression) may inhibit ULK1 phosphorylation but may not contribute to uncoupling between AMPK and ULK1.

Interestingly, ectopic expression of acid ceramidase restored ULK1 phosphorylation in a nutrient-laden environment, but it did not restore LC3 protein levels (data not shown), alter the effects of AMPK activators on LC3 (data not shown), nor reduce accumulation of p62 protein in nutrient-laden HAECs (data not shown). This indicates that while ceramides may be important regulators of ULK1, the latter is not the only contributor to autophagy regulation under these conditions.

**DISCUSSION**

Regulation of AMPK plays a key role in events associated with type 2 diabetes and the metabolic syndrome, as well as their associated cardiovascular complications (24, 87). For example, two of the most widely prescribed drugs for diabetes and cardiovascular disease, metformin and statins, are both
AMPK activators (82, 95). While these treatments significantly help control diabetes and reduce cardiovascular risk across the population, they are not effective in all patients (10, 37, 79). This suggests that there may be certain pathological conditions in which AMPK becomes uncoupled from its expected cellular actions. Under these conditions (such as the model described here), there may be a need for additional therapeutic targets independent of AMPK.

In this study, we treated primary HAECs with a combination of high glucose and palmitate to model the pathological stress of poorly controlled type 2 diabetes. In this context, we analyzed the role of AMPK in regulation of autophagy, a pathological process implicated in atherogenesis. Consistent with the role of these nutrients in endothelial dysfunction (22, 47) as well as previous studies in HUVECs (8, 9, 41), we found that the combination of glucose and palmitate both predisposed HAECs to apoptosis and increased their inflammatory profile. The subsequent novel findings from our work were as follows: 1) Incubation of HAECs with the same high concentrations of glucose and palmitate that increased apoptosis and inflammation also impaired autophagy. 2) Exposure of HAECs to excess nutrients decreased the activity of AMPK. 3) Unlike low nutrient conditions, chemical activation of AMPK did not restore autophagy in this nutrient-laden environment. 4) Ectopic expression of acid ceramidase prevented nutrient-induced decreases inULK1 phosphorylation at Ser555, an AMPK-targeted site. Surprisingly, increased expression of acid ceramidase did not alter the effects of AICAR, A769662, or phenformin on ULK1 phosphorylation. This suggests that accumulation of ceramides may directly regulate ULK1, but perhaps not through interfering with the actions of AMPK. Collectively, these data indicate that, in HAECs, high concentrations of glucose and palmitate exert a metabolic stress (perhaps similar to those of patients who do not respond to AMPK activators) under which AMPK activation does not induce autophagy. Intriguingly, restoration of ULK1 phosphorylation by acid ceramidase expression suggests that targeting ceramide or its metabolites may be a potential alternative target to allay nutrient-induced stress in the endothelium.

The complex interactions between ULK1, AMPK, and mTORC1 that regulate starvation-induced autophagy are incompletely understood, but even less is known about their relationships when there is an excess of nutrients (13, 63). The inability to restore autophagic flux in HAECs by chemically activating AMPK contrasts with previous observations in rodent myocardial cells (34) and macrophages (88). He and colleagues (34) observed that hyperglycemia, induced by streptozotocin or incubating myoblasts in a high-glucose media (30 mM for 48 h), suppressed autophagy; likewise Wen and colleagues found that 24 h of exposure to 0.5 mM palmitate, following stimulation with lipopolysaccharide, suppressed autophagy in rodent macrophages (88). In each of these contexts, autophagy impairment induced by a single nutrient was overcome by AMPK activation. AMPK activation in our human endothelial cell model may have been insufficient to rescue nutrient-induced loss of autophagy due to the presence of a double nutrient (glucose with palmitate), rather than a single nutrient (glucose or palmitate)-induced inhibition of flux. The increased severity of a double nutrient—compared with a single nutrient—stress is supported by data from the β-cell, in which a combined exposure to glucose and palmitate produced a more severe inhibition of autophagic flux than exposure to palmitate alone (58). In addition to the presence of a double nutrient stress, our model differs from the above mentioned studies in its use of a chronic glucose treatment. Exposure of HAECs to high concentrations of glucose during cell division and throughout the lifetime of the newly generated cells may alter their epigenetic profile, as shown in other types of cells (67), thus altering their responses to chemical activators such as AICAR.

Given that acid ceramidase expression did not facilitate additional ULK1 phosphorylation or autophagosome formation with AICAR or A769662, the mechanism by which glucose and palmitate alter the interaction between AMPK and autophagy remains an open area of investigation. There may be separate pools of AMPK in which signaling to its respective targets (ACC or ULK1) is differentially regulated. This could be a result of glucose- and palmitate-induced posttranslational modifications on AMPK and/or ULK1 that prevent interactions between these two molecules. Interestingly, in the presence of excess nutrients, acid ceramidase expression restored phosphorylation of ULK1 (Fig. 6C and data not shown) and ACC, but not AMPK (data not shown), suggesting that acid ceramidase may be involved in both of these signaling pathways, perhaps independent of AMPK. There are several potential mechanisms by which ceramides or other sphingolipid metabolites may be impairing autophagy, including activation of protein phosphatases (65, 85) and increased levels of endoplasmic reticulum stress (5, 28, 61). Further study is needed to focus on the contribution of these lipids, as well as other processes such as the inhibition of Sirtuin-1 (SIRT1) (31, 53, 59, 83) or generation of oxidative stress (78) to nutrient-induced autophagy impairment.

In addition to modulating the initiation stages of autophagy through ULK1, excess nutrients may also affect initiation and autophagosomal membrane elongation through inhibition of SIRT1, an NAD⁺-dependent histone deacetylase. We and others have shown that high concentrations of glucose or palmitate impair AMPK activity (35, 36, 91) as well as SIRT1 activity (2, 18, 71, 80). AMPK and SIRT1 have also been shown to regulate one another (11, 39, 55). In the nucleus, SIRT1 may activate autophagy by deacetylating several members of the forkhead family of transcription factors, FOXO1 and FOXO3 (31, 53). In the cytosol, in vitro studies indicate that Atg5, Atg7, and Atg8, autophagy proteins required for the formation and elongation of the autophagosomal membrane (46), are also targets of SIRT1 (59). In mouse embryonic fibroblasts, overexpression of wild-type SIRT1, but not deacetylase-deficient SIRT1, increased basal autophagy (59). Consistent with this autophagy-activating characteristic of SIRT1, inhibition of SIRT1 in human THP-1 cells impaired autophagy (83).

Perhaps related to SIRT1 inhibition, excess concentrations of glucose and palmitate may also impair autophagy through the generation of oxidative stress (4, 8, 15, 40, 72). Reactive oxygen species inhibit the activity of Atg4 (78), an enzyme required both for the maturation of LC3 prior to its incorporation into the forming autophagosomal membrane and for LC3 cleavage from the outside of the autolysosome to be recycled for future use (46). Thus, inhibition of Atg4 via
reactive oxygen species may impair the incorporation of both newly synthesized and recycled LC3 into autophagosomes. LC3 is critical for the elongation of the autophagosomal membrane and depletion of LC3 has been associated with smaller autophagosomes (92), which would consequently degrade fewer substrates.

In addition to decreasing autophagosome size, glucose and palmitate may also impair substrate degradation in the autolysosome by interfering with lysosomal proteases, such as cathepsin L (Fig. 2D). Evidence from pancreatic β-cells and hepatocytes suggests that nutrient-induced loss of degradative capacity may be due to inhibition of lysosomal acidification (58) or increased lysosomal permeabilization (25).

In summary, we have identified autophagic dysregulation as a new mechanism linking glucose and palmitate exposure with inflammation and apoptosis in HAECs. Our HAEC model of poorly controlled type 2 diabetes, which utilizes chronic exposure to glucose in combination with acute palmitate treatment, revealed that these nutrients impair ULK1 phosphorylation, autophagosome formation, and cathepsin L activity (Fig. 7). Similar to the effects of glucose and palmitate on AMPK activity in rat adipocytes (35), cardiomyocytes (36), and bovine aortic endothelial cells (91), we found that these nutrients decreased AMPK activity in HAECs. However, activation of AMPK was not sufficient to restore autophagy in this nutrient-rich environment. Thus, unlike the role ascribed to AMPK in starvation-induced autophagy, AMPK is uncoupled from autophagy under nutrient excess conditions. Rather than exclusively targeting AMPK, ceramide or its metabolites may be additional targets for therapies aimed at preventing autophagic dysregulation in excess nutrient conditions. Moreover, an improved understanding of the particular types and magnitudes of stress under which AMPK activation can restore cellular homeostasis and those in which it cannot, may lay the foundation for improved treatments for a variety of metabolic pathologies in which AMPK plays a critical role.

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**DISCLOSURES**

No conflicts of interest, financial or otherwise, are declared by the author(s).

**AUTHOR CONTRIBUTIONS**

K.A.W. and Y.I. conception and design of research; K.A.W. and Y.I. performed experiments; K.A.W. and Y.I. analyzed data; K.A.W. and Y.I. interpreted results of experiments; K.A.W. prepared figures; K.A.W. drafted...
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