Mangiferin Stimulates Carbohydrate Oxidation and Protects Against Metabolic Disorders Induced by High-Fat Diets

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Excessive dietary fat intake causes systemic metabolic toxicity, manifested in weight gain, hyperglycemia, and insulin resistance. In addition, carbohydrate utilization as a fuel is substantially inhibited. Correction or reversal of these effects during high-fat diet (HFD) intake is of exceptional interest in light of widespread occurrence of diet-associated metabolic disorders in global human populations. Here we report that mangiferin (MGF), a natural compound (the predominant constituent of Mangifera indica extract from the plant that produces mango), protected against HFD-induced weight gain, increased aerobic mitochondrial capacity and thermogenesis, and improved glucose and insulin profiles. To obtain mechanistic insight into the basis for these effects, we determined that mice exposed to an HFD combined with MGF exhibited a substantial shift in respiratory quotient from fatty acid toward carbohydrate utilization. MGF treatment significantly increased glucose oxidation in muscle of HFD-fed mice without changing fatty acid oxidation. These results indicate that MGF redirects fuel utilization toward carbohydrates. In cultured C2C12 myotubes, MGF increased glucose and pyruvate oxidation and ATP production without affecting fatty acid oxidation, confirming in vivo and ex vivo effects. Furthermore, MGF inhibited anaerobic metabolism of pyruvate to lactate but enhanced pyruvate oxidation. A key target of MGF appears to be pyruvate dehydrogenase, determined to be activated by MGF in a variety of assays. These findings underscore the therapeutic potential of activation of carbohydrate utilization in correction of metabolic syndrome and highlight the potential of MGF to serve as a model compound that can elicit fuel-switching effects.

The worldwide incidence of diabetes was estimated to be 347 million in 2008 and has been projected to continue to increase (1). The most important driving force for this high incidence of diabetes is an increasing prevalence of obesity (2) linked to consumption of high-calorie foods and sedentary activity. Although lifestyle changes can mitigate some of the prevailing metabolic syndrome in obese populations, targeted therapies that modify fundamental cellular and physiologic processes are needed for long-term medical success in treating this epidemic. Cellular dysfunction includes mitochondrial deterioration, which has been implicated as a fundamental feature accompanying type 2 diabetes. Previous studies have demonstrated that pharmacologic or genetic upregulation of mitochondrial functions can provide favorable outcomes in diet-induced metabolic syndromes (3,4).

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Although enhancing fatty acid oxidation can reduce fat accumulation, it may not directly mitigate hyperglycemia, because fatty acid oxidation inhibits glucose utilization (5,6). Dietary overconsumption of fats/lipids enhances fatty acid fuel utilization but induces metabolic inflexibility and insulin resistance (7), even when accompanied by increased mitochondrial density and increased oxidative capacities (8,9). This conundrum at least theoretically weakens strategies for increasing fatty acid oxidation as a means to rescue animals subject to metabolic disorders induced by a high-fat diet (HFD). We argue that an alternative and key consideration associated with strategies that seek enhanced oxidative metabolism is whether they can promote glucose utilization, which could improve glucose profiles and mitigate insulin resistance.

Skeletal muscle is the major peripheral tissue that removes blood glucose, accounting for ~75% of whole-body glucose uptake (6). After uptake into tissues, glucose is metabolized into different pathways, with a major catabolite being pyruvate, a central branch-point metabolite in carbohydrate metabolism. Subject to regulatory factors, pyruvate is converted to several downstream metabolites, including alanine, oxaloacetate, lactate, and acetyl-CoA, with the last two as major metabolites (10). The irreversible conversion of pyruvate to acetyl-CoA, catalyzed by pyruvate dehydrogenase (PDH), provides a regulatory step by which glucose enters into oxidative metabolism. Alternatively, conversion of pyruvate to lactate by lactate dehydrogenase (LDH) (10) enables muscle release of lactate for the Cori cycle from muscle to liver.

PDH serves as a central gatekeeper of carbohydrate oxidative metabolism. PDH activity regulates the fate of carbohydrate metabolism in muscle as the determining factor in setting the competition between glucose and fatty acid catabolism in this tissue (7,11). Decreased PDH activity is proposed to be a fundamental factor in the derangement of carbohydrate metabolism (12) observed in diabetic heart and skeletal muscle (13,14). Activation of PDH by dichloroacetate (DCA) or by deletion of the negative regulator, PDH kinase (PDK), improves muscle functions, lowers glucose levels, and increases glucose tolerance and insulin sensitivity in mice and humans (15,16). These data indicate that improved carbohydrate utilization linked to activation of key metabolic steps can provide fundamental improvements in metabolic syndrome.

In this study, we report that stimulation of carbohydrate oxidation is the pharmacodynamic outcome of a natural product, mangiferin (MGF), the predominant constituent of Mangifera indica bark extract (mango tree; Supplementary Fig. 1). Mice treated with MGF were protected from HFD-induced toxicities, including weight gain, insulin resistance, and glucose intolerance. Further investigation into its mechanisms of actions demonstrated that MGF activates PDH, thereby increasing carbohydrate oxidation and metabolic flexibility of energy substrate utilization.

**RESEARCH DESIGN AND METHODS**

Chemical reagents were purchased from Sigma-Aldrich (St. Louis, MO), unless specified. Animals were fed a chow diet (CD; Purina LabDiet, Framingham, MA) providing 28.5% (kcal) protein, 13.5% fat, and 58% carbohydrates, or an HFD (Research Diets, New Brunswick, NJ) providing 20% (kcal) protein, 60% fat, and 20% carbohydrates.

**Animal Experiments**

Wild-type C57BL6/J mice at the age of 5–6 weeks were purchased from The Jackson Laboratory (Bar Harbor, ME). All studies were approved by and performed in compliance with the guidelines of the Yeshiva University Institutional Animal Care and Use Committee. The CD and HFD were ground into fine powder and mixed with pure MGF (CTMedChem, Bronx, NY). The mixtures were repelled and administered to individually housed mice. Mice were fed ad libitum with the CD or HFD, with or without MGF, and water.

Fat composition and fat distribution were determined by MRI. Lean and fat mass was determined as described (17). Metabolic measurements (V̇O₂), respiratory quotient (RQ), energy expenditure (EE), and locomotor activity were obtained continuously using a CLAMS (Columbus Instruments) open-circuit indirect calorimetry system (18). Animals were acclimatized for 2 days, and data were subsequently collected over a 5-day period. Body temperature was measured using a BAT-10 thermometer (Physi temp, Clifton, NJ). Hyperinsulinemic-euglycemic clamps were performed as described (19). An intraperitoneal glucose tolerance test (IGTT) was performed on conscious mice after 12-h fasting and an intraperitoneal injection with 20% glucose in saline (2 g/kg total body weight).

**Fatty Acid and Glucose Oxidation in Isolated Muscle**

Palmitate and glucose oxidation in soleus muscle were measured using previously described methods (20,21), with slight modification. Soleus muscle was incubated in the oxidation buffer for 1 h for palmitate oxidation and for 6 h for glucose oxidation.

**Cell Culture**

C2C12 myoblast cells were maintained in DMEM high glucose (HyClone Laboratories, Logan, UT), supplemented with 1% penicillin and streptomycin (Gibco, Carlsbad, CA) and 10% FBS. C2C12 cells were differentiated in the above DMEM medium containing 2% horse serum (replacing 10% FBS) for 4–6 days after cells reached confluence.

Fatty acid oxidation in myotubes was measured by radioactivity assay using methods as described (22).

**Oxygen Consumption Rate and Extracellular Acidification Rate Measurements by Metabolic Flux Analyzer**

Oxygen consumption rate (OCR) and extracellular acidification rate (ECAR) in C2C12 myotubes were obtained by a Metabolic Flux analyzer (Seahorse Bioscience), as previously described (23,24). In brief, myotubes differentiated...
from C2C12 cells in XF24 microplates were incubated overnight in glucose-free and sodium pyruvate-free DMEM containing l-glutamine, 2% horse serum, and 1% penicillin/streptomycin, with or without 200 μmol/L MGF. The concentrations of MGF used for this and the subsequent in vitro assays were chosen based on the in vivo effective dose of 400 mg/kg body weight and oral MGF bioavailability of 1.2% in rat (25). Medium containing vehicle, BSA-conjugated palmitate, d-glucose, or sodium pyruvate was loaded in port A of the sensor cartridge, and 70 μL was injected to wells containing 630 μL media. The final concentrations of BSA-palmitate, d-glucose, and sodium pyruvate were 0.1, 25, and 50 mmol/L, respectively.

**Mitochondrial Protein Preparation**

Mitochondria were isolated from C2C12 myotubes treated with various concentrations of MGF for 24 h or with various concentrations of MGF, 5 mmol/L DCA, or 2 mmol/L fluoropyruvate (FP) plus 0.2 mmol/L MGF for 15 min, at 37°C, using the previously described method (26,27) with modified buffer (250 mmol/L sucrose, 1 mmol/L EGTA, 10 mmol/L HEPES, and 5 g/L BSA, pH 7.5). Mitochondrial pellets were suspended, and mitochondrial protein was extracted by a freeze-thaw procedure (26,27).

**PDH Activity**

**Spectrometry Assay**

As adapted from previous reports (26,28,29), 100 μg mitochondrial protein lysate was loaded in a 96-well plate in 200 μL reaction buffer containing 5 mmol/L l-carnitine, 2.5 mmol/L NaOH. A 25 mmol/L NAD+, 0.2 mmol/L thiamine pyrophosphate, 0.1 mmol/L CoA, 1 mL/L Triton X-100, 0.3 mmol/L DTT, 1 mmol/L MgCl2, 1 g/L BSA, and 6.5 mmol/L phenazine methosulfate in 50 mmol/L potassium phosphate buffer (pH 7.5). Reaction was initiated by adding 5 mmol/L pyruvate (50 μL) and 0.6 mmol/L p-iodonitrotetrazolium violet.

**Radioactivity Assay**

We developed a system based on previous reports (27,30–32), in which a manifold was used to split the airflow into six branches that were simultaneously connected to six pairs of reaction and trapping vials. Into each reaction vial was added 200 μL reaction buffer (1 mmol/L MgCl2, 1 mmol/L DTT, 0.1 mmol/L thiamine pyrophosphate, 0.3 mmol/L CoA, 2.5 mmol/L NAD+, 50 mmol/L potassium phosphate, 75 μmol/L sodium pyruvate, freshly prepared, pH 8.0), and 0.2 μL [carboxy-14C]-labeled pyruvate (0.1 μCi/μL, 55 μCi/μmol). Into each trapping vial was added 300 μL trapping solution (1 mmol/L NaOH). After airflow was stabilized, mitochondrial protein lysate (10–50 μg) was injected into the reaction vial. At various time points (3–40 min), reactions were stopped by injecting 10% trichloroacetic acid (250 μL), and 14CO2 was trapped for 3 h. Reaction and trapping were performed at room temperature. 14C in trapping and reaction vials was counted separately by a scintillation counter. Statistical analyses are specified in the figure legends.

**RESULTS**

**MFG Protects Against HFD-Induced Obesity**

A report indicated that MGF treatment in a type 2 diabetic mouse model (KK-Ay mice) improved the glucose and insulin profile (33). To evaluate MGF effects on HFD-induced insulin resistance and obesity, C57BL/6J mice were fed the CD or HFD containing 0%, 2.5%, or 5.0% g MGF/g food. Mice fed the CD supplemented with MGF exhibited similar weight gain as controls (Fig. 1A). When challenged with the HFD, mice experienced substantial weight gain compared with CD mice over a 16-week period. MGF treatments decreased HFD-induced weight gain in a dose-dependent manner (Fig. 1A). After 5 weeks, 0.5% MGF (~400 mg/kg as a result of average daily food intake of 2–3 g) produced statistically significant weight reduction. Impressively, continued treatment with MGF

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**Figure 1** — MGF mitigates HFD-induced obesity. C57BL6J mice (5–6 weeks old) were fed with CD or HFD, with or without MGF (0, 0.25%, 0.5% g/g), for 16 weeks (n = 10 per group). Body weight was monitored daily. A: Graph of body weight gain over time. *P < 0.05 indicates significant changes between HFD and HFD + 0.5% MGF groups. B: Body composition evaluated by nuclear magnetic resonance after 16 weeks of diet. *P < 0.05. Values are average ± SEM. C: MRI of two representative mice fed with HFD or HFD + 0.5% MGF for 16 weeks. “Water on resonance” is the typical MRI image at the level of the kidneys. “Fat on resonance” shows the same anatomic location with the transmitter set on the fat resonance to show fat distribution. P values were obtained by ANOVA test.
for 16 weeks prevented 50% of the body weight gain of the HFD-fed mice (Fig. 1A). Nuclear magnetic resonance–based whole-body fat quantitation revealed that MGF decreased total fat mass (Fig. 1B), accounting for most of the weight difference observed between treated and untreated animals. MGF also modestly reduced lean mass (Fig. 1B) because lean mass loss often accompanies fat mass loss (34). MRI cross-sectional images showed that abdominal fat accumulation was largely prevented by MGF treatment (Fig. 1C).

**MGF Enhances Systemic Respiration and EE**

To further investigate the physiologic basis for the reduced body weight gain in MGF-treated animals, we quantified caloric intake and EE by indirect calorimetry (Fig. 2). MGF did not affect food or water intake (Fig. 2A). In CD-fed mice, MGF treatment caused a slight increase in activity in the dark cycle (Fig. 2B) but no change in total spontaneous locomotor activity (Fig. 2C). In HFD-fed mice, MGF modestly increased locomotor activity in nocturnal hours. EE was increased in mice fed the MGF-treated HFD (Fig. 2D and E), which was correlated with increased $V_O^2$ (Supplementary Fig. 2). The increase in spontaneous locomotor activity was marginal and consistent with spontaneous locomotor activity being insufficient to account for alterations in EE in standard mouse caging (35), suggesting that MGF-caused increases in EE were from activity-independent metabolic processes.

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**Figure 2**—MGF enhances EE and promotes carbohydrate utilization in vivo. C57BL6/J mice (5–6 weeks old) were fed with CD or HFD or those diets mixed a dose of 0.5% MGF for 16 weeks ($n=10$ per group). A: Average daily food and water intake. Indirect calorimetry assessments of locomotor activity (B and C), EE (D and E), and RQ (G and H). F: Cold test. Body temperature was monitored while mice were exposed to 4°C for 5 h. *$P < 0.05$. $P$ values were obtained by t test for F and by ANOVA test for the rest. Values are average ± SEM.
An additional pathway for dissipation of energy is thermogenesis. MGF modestly increased basal body temperature when mice were maintained at an ambient temperature (Fig. 2F). This effect was more pronounced when mice were exposed to cold, highlighting an increased thermogenic capacity in MGF-treated animals, probably resulting from MGF-induced uncoupling proteins in brown adipose tissue (Supplementary Fig. 3).

**MGF Stimulates Carbohydrate Utilization and Increases Flexibility in Use of Energy Substrates**

We measured RQ to investigate whether MGF exerted changes in energy metabolism and metabolic fuel source selection. In untreated CD mice, RQ was ~0.9 during light hours and gradually increased during dark hours to 0.93, and MGF had a minimal effect (Fig. 2G and H). In untreated HFD-fed mice, RQ averaged at 0.77 at all times, demonstrating a striking shift to fatty acids as preferred fuels and apparent metabolic inflexibility (Fig. 2G and H). MGF treatment in HFD-fed mice increased RQ and RQ fluctuation, causing a range of 0.81–0.85 during the light and dark cycles (Fig. 2G and H). These data indicate that MGF promotes carbohydrate oxidation and improves fuel source flexibility.

**MGF Protects Against HFD-Induced Glucose Intolerance and Insulin Resistance**

Fasting plasma glucose was increased to 198 ± 25 mg/dL in 18-week HFD-fed mice compared with 116 ± 16 mg/dL in CD-fed controls (Fig. 3A). MGF treatment had no significant effect on glucose levels in CD mice but reduced the fasting glucose levels of HFD mice to 132 ± 7 mg/dL. HFD feeding drastically increased plasma insulin level to 2.93 ± 0.48 ng/mL from ~0.75 ± 0.12 ng/mL in CD mice. MGF treatment resulted in a complete normalization of insulin levels in the HFD mice (Fig. 3B). To directly assess insulin sensitivity, we performed hyperinsulinemic-euglycemic clamp studies. As expected, HFD animals were insulin resistant compared with controls, as evidenced by a lower glucose infusion rate (GIR; Fig. 3C). MGF tripled the GIR in HFD animals, indicating that MGF improves insulin sensitivity.

One of the key factors affecting the GIR is the rate of glucose disposal by peripheral tissues such as muscle. MGF tripled the glucose disposal rate in HFD mice (Fig. 3D), providing direct evidence that MGF stimulates glucose utilization in muscle during HFD feeding. In an IGTT, MGF-treated HFD mice exhibited significantly improved glucose tolerance versus HFD mice, which were markedly glucose intolerant compared with CD mice (Fig. 3E). MGF-treated animals exhibited lower insulin levels at basal conditions and during the IGTT (Fig. 3F), consistent with MGF improvement of insulin sensitivity. MGF also modestly but significantly increased total plasma adiponectin in HFD mice (Fig. 3G) and prevented HFD-induced increase in plasma triglycerides (Fig. 3H), without any significant effect on free fatty acids (Fig. 3I).

**MGF Promotes Carbohydrate Utilization**

Improved glycemia and insulin sensitivity led us to connect the mechanism of action of MGF promotion of carbohydrate oxidation as indicated by the RQ. Glucose uptake data from the hyperinsulinemic-euglycemic clamp (Fig. 3D) also indicated enhanced glucose disposal and increased glucose catabolism. To further investigate these effects, we measured palmitate and glucose oxidation in the soleus skeletal muscle. HFD feeding doubled palmitate oxidation and conversely reduced glucose oxidation, consistent with prior reports (20,36). Interestingly, MGF did not significantly affect palmitate oxidation (Fig. 4A) but significantly increased glucose oxidation by 60% in HFD mice (Fig. 4B), suggesting that MGF preferentially enhances glucose utilization in the muscle.

To further evaluate MGF effects on fuel preference, fatty acid and glucose oxidation were both measured in cultured C2C12-derived myotubes. MGF did not significantly affect [carbonyl-13C]oleic acid oxidation in myotubes as measured by a radioactivity assay (Fig. 4C) nor did it affect palmitate OCR, as measured by the Seahorse Bioscience Metabolic Flux analyzer (Fig. 4D), consistent with the ex vivo results (Fig. 4A). We then monitored glucose catabolism in C2C12 myotubes by the Metabolic Flux analyzer. When treated with MGF, the myotube glucose oxidation rate was increased by ~50% (Fig. 4E). In addition, MGF accelerated glucose utilization as measured by time to peak (Fig. 4E). Some of this effect may be linked to MGF increased GLUT4 mRNA in the soleus muscle of HFD-fed mice and in C2C12 cells (Supplementary Fig. 4). These observations are consistent with ex vivo and in vivo results, confirming that MGF increases glucose metabolism, independent of fatty acid oxidation, even under a high dietary fat challenge. The MGF-stimulated increase in glucose metabolism also resulted in a statistically significant and dose-dependent increase in ATP production (Fig. 4F).

**MGF Inhibits Lactic Acidosis**

The end metabolite of glycolysis is pyruvate, which is a key intersection metabolite that can be converted by LDH to lactate or catabolized by PDH to acetyl-CoA. We first investigated whether MGF affected lactate formation. Using the Seahorse Bioscience analyzer, lactate production was monitored by the extracellular acidification of the media, ECAR of the media (23,24). Myotubes treated with glucose exhibited marked acidification of media, as measured by ECAR. Interestingly, MGF significantly blunted this effect (Fig. 5A).

Because ECAR is an indirect measure of lactate production, we directly measured lactate in the media in the Seahorse Bioscience plate. Indeed, media lactate in glucose-treated myotubes accumulated to ~80 nmol/L, and MGF reduced lactate accumulation to 30 nmol/L (Fig. 5B). In separate experiments, we determined that MGF inhibited lactate accumulation when glucose was fed to cells in a dose-dependent manner, with a half-maximal inhibitory concentration of 337 ± 40 µmol/L (Fig. 5C), demonstrating...
that MGF indeed inhibits lactate formation from glucose. Reduced lactate formation could be caused by direct inhibition of LDH or reduced availability of pyruvate to the fermentation pathway. We measured LDH activity using a kinetic assay with and without MGF and noted no effect on LDH activity (data not shown), suggesting that MGF might affect lactate production by enhancing pyruvate oxidation through PDH.

**MGF Enhances Pyruvate Oxidation by Activating PDH**

To determine if MGF activates PDH, C2C12 myotubes were incubated with pyruvate and displayed increased OCR. The addition of MGF with pyruvate resulted in both a larger OCR and temporal rate of activation (Fig. 6A), verifying that MGF enhances and accelerates oxidation of pyruvate (Fig. 6A) and glucose (Fig. 4E) as fuel substrates,
but not fats (Fig. 4C and D), in myotubes. These data imply that metabolic mediators downstream of pyruvate could be pharmacodynamic targets of MGF. We therefore investigated a putative target, PDH, using two enzymatic activity assays. One is a spectrometry assay measuring the pyruvate-dependent production of NADH (28,29,37), and the other is a radioactivity assay directly measuring $^{14}$CO$_2$ production from [carboxy-$^{14}$C]pyruvate oxidation. Using the spectrometry assay, we observed that MGF increased NADH production in a dose-dependent manner and was more effective than the known PDH activator DCA (30) (Fig. 6B). To confirm that the NADH increase was PDH-dependent, we added FP, a potent inhibitor of PDH. Upon addition of FP, MGF stimulation of NADH production was abolished, suggesting that MGF acts to increase NADH production via increases in PDH activity. From the absorbance versus time plots, we analyzed initial velocities of NADH formation and determined that MGF dose-dependently increased the initial rates of PDH-catalyzed reactions (Fig. 6C). Treatments for 24 h caused greater increases than 15-min treatments, although the latter contributed a large portion of the former. The half-maximal effective concentrations (EC$_{50}$) of 24-h and 15-min MGF treatments were $184 \pm 23$ and $216 \pm 35$ μmol/L, respectively. This suggests that effects of MGF on PDH may be complex, having acute as well as inducible features.

To confirm these studies, we sought a more reliable assay that can directly measure PDH activity. Radiolabeled [carboxy-$^{14}$C]pyruvate substrate was used to measure PDH activity. We determined that acute treatment with MGF for 15 min provides statistically significant
increases in PDH activity, with an EC$_{50}$ of 234 ± 37 µmol/L (Fig. 6D). The two sets of data obtained from two distinct methods (Fig. 6C, filled symbols; Fig. 6D) are in agreement, consistently showing that apparent PDH activity increased several fold and was ultimately saturated.

PDH is regulated by reversible phosphorylation by PDK(s) and PDH phosphatases (38). MGF treatment for 24 h resulted in a dose-dependent decrease in phosphorylated PDH (p-PDH) (Fig. 6E and F), consistent with increased PDH activity. Among four PDKs, PDK4 is highly expressed in muscle and sensitive to nutritional regulation and diabetic state (39,40). We assessed the effects of MGF on PDK4 and found that MGF dose-dependently suppressed PDK4 protein expression (Fig. 6E and F). In cells acutely treated with MGF for 15 min, MGF also reduced p-PDH (Fig. 6G and H), but did not change PDK4 expression level (data not shown), suggesting that acutely reduced p-PDH by MGF is probably due to inhibition of PDK activity. Inhibition of PDK(s) is the mechanism by which DCA activates PDH. MGF is more effective than DCA (Fig. 6G and H).

**DISCUSSION**

Excess dietary fat consumption and consequent excessive fatty acid utilization cause toxic impairments to glucose utilization, resulting in hyperglycemia, insulin resistance, and weight gain. In this respect, poor glucose utilization by the skeletal muscle has been identified as a major contributor to insulin resistance and weight gain associated type 2 diabetes (7). Strategies aimed to stimulate skeletal muscle bioenergetics as a direct approach to counteract diabetes have been pursued but have focused principally on mitochondrial activation and, particularly, increasing fatty acids oxidation. This strategy has been demonstrated to be of therapeutic interest and has some impressive successes, but does have limitations because it does not specifically address poor carbohydrate utilization and possible corrective effects obtainable by increasing carbohydrate utilization as opposed to increasing fatty acid oxidation.

Here we report novel findings that address the fundamental question of whether it is feasible to correct metabolic syndrome by improving carbohydrate utilization selectively versus fatty acids. We discovered that MGF stimulates carbohydrate utilization even when fat is the predominant energy source. MGF was able to mitigate the negative consequences of excessive fat consumption and protect against diet-induced weight gain, hyperglycemia, glucose intolerance, hyperinsulinemia, and insulin resistance (Figs. 1 and 3A–F). In addition, it increases EE and preserves fuel utilization flexibility (Fig. 2D, E, G, and H). Increased RQ values with MGF treatment are consistent with stimulated carbohydrate utilization (Fig. 2G), which was confirmed in isolated muscle tissues and in cultured myotubes (Fig. 4).

Furthermore, we investigated the effects of MGF on the central metabolic switcher, PDH. The data from two assays consistently demonstrate that long- and short-term MGF treatments both potently increase PDH activity (Fig. 6B–D) and that this increase is, at least partially, due to reduced p-PDH resulting from MGF downregulation of PDK4 (Fig. 6E–H). HFD induces PDK4 (27,41), a mechanism by which HFD inhibits PDH activity, resulting in inhibition of carbohydrate utilization (41). MGF is able to suppress PDK4, and understandably, this effect could be more profound when PDK4 is induced by an HFD, which explains why the MGF-caused glucose oxidation increase under an HFD is greater than under a CD (Fig. 4B).

In line with the Randle cycle hypothesis (5,7), although MGF significantly enhanced carbohydrate oxidation, there
Figure 6—MGF enhances pyruvate oxidation by activating PDH. A: Pyruvate oxidation in myotubes differentiated from C2C12 cells measured by Seahorse Bioscience Metabolic Flux analysis. Myotubes differentiated from C2C12 cells in XF24 microplates were incubated overnight in glucose-free and sodium pyruvate–free DMEM, with or without 200 μmol/L MGF. OCR was recorded after injection of sodium pyruvate into wells, with or without 200 μmol/L MGF. B: PDH activities measured as the difference between absorbances at 500 nm and 750 nm at 25°C in mitochondrial protein from C2C12 myotubes treated with various concentrations of MGF for 24 h or 15 min or with 5 mmol/L DCA or FP + 200 μmol/L MGF for 15 min. C: Initial rates of PDH-catalyzed pyruvate oxidation measured as the difference between absorbances at 500 nm and 750 nm in mitochondrial protein from C2C12 myotubes treated with various concentrations of MGF for 24 h or 15 min. The unit of activity was determined by using extinction coefficient of reduced p-iodonitrotetrazolium violet (12.4 mmol·L⁻¹·cm⁻¹). D: Initial rates of PDH catalyzed [¹⁴C]-labeled pyruvate oxidation measured as the production of ¹⁴CO₂ by mitochondrial protein from C2C12 myotubes treated with various concentrations of MGF for 15 min. E: Western blots of proteins extracted from C2C12 myotubes treated with 200 or 1,000 μmol/L MGF for 15 min and 24 h. Membrane was blotted with PDH (1:250 dilution; Cell Signaling), p-PDH (1:2,000; Abcam), PDK4 (1:500; laboratory of Dr. Robert Harris at Indiana University School of Medicine), or α-tubulin (1:2,000; Sigma-Aldrich) at 4°C overnight. F: Quantitative analysis of Western blots shown in E. G: Western blots of proteins extracted from C2C12 myotubes treated with 200 or 1,000 μmol/L MGF or 5 mmol/L DCA for 15 min. H: Quantitative analysis of Western blots shown in G. Values are average ± SEM. *P < 0.05. P values were obtained by t test (A) or ANOVA test (F and H).
was a trend for reduced fatty acid oxidation in both muscle and C2C12 myotubes (Fig. 4A, C, and D). In hepatic tissues and cells, Niu et al. (42) observed that MGF increased oleic acid uptake and induced carnitine palmitoyltransferase 1, suggesting that MGF possibly enhances fatty acid utilization in liver. These differences could reflect tissue-specific effects of MGF.

The PDH-catalyzed product acetyl-CoA can exit mitochondria to yield cytoplasmic acetyl-CoA, which serves as the substrate for de novo lipogenesis, suggesting that activation of PDH by MGF could potentially increase lipogenesis. Interestingly, our unbiased proteomics study revealed that MGF downregulated several critical enzymes in lipogenesis and predicted that MGF suppressed SREBP-1 (43), an important regulator of lipogenesis and adipogenesis (44,45). Further investigation uncovered that MGF inhibited adipogenesis in 3T3 L1 cells, probably by downregulating SREBP, as well as C/EBP and peroxisome proliferator–activated receptor-γ (data not shown). These inhibitory effects of MGF on lipogenesis and adipogenesis could be potential mechanisms by which MGF prevents HFD-induced obesity.

In addition, MGF enhanced adaptive thermogenesis (Fig. 2F), likely by activating brown adipose tissue uncoupling mechanisms (Supplementary Fig. 3). Besides inducing uncoupling proteins and presumably increasing energy leak/waste, MGF increases energy production as it increases ATP levels (Fig. 4F and Supplementary Fig. 5). We have shown that MGF induces enzymes catalyzing the tricarboxylic acid cycle and the electron transport chain (43). In addition, MGF raises the NAD⁺-to-NADH ratio (Supplementary Fig. 6), which propels the flux through the tricarboxylic acid cycle and the electron transport chain. These data suggest that MGF enhances mitochondrial bioenergetics, supporting the previous findings that MGF activates AMPK (42) and that activation of AMPK raises the NAD⁺-to-NADH ratio and increases mitochondrial bioenergetics (46).

Enhancing the mitochondrial respiratory chain could potentially increase reactive oxygen species. Interestingly MGF has well-established antioxidant properties because it bears a catechol moiety that enables it to form stable MGF-Fe²⁺/Fe³⁺ complexes, preventing Fenton-type reaction and lipid peroxidation (47). This antioxidant property could facilitate and maintain MGF activation of PDH, because PDH is susceptible to inactivation by reactive oxygen species (48). In turn, activation of PDH prevents O₂⁻ accumulation because PDH deficiency is associated with impaired mitochondrial O₂⁻ removal (49).

In summary, we report a novel small molecule, MGF, that is able to stimulate carbohydrate oxidation when fatty acid is the dominant energy source by enhancing the activity of PDH, a critical metabolic switcher, in the regulation of overall energy balance and homeostasis. Owing to its chemical structural features, oral MGF is bioavailable in rodent and human (25,50). Although the bioavailability of oral MGF is low, demanding a high effective dose in vivo, and the EC₅₀ of MGF is in the micromole/liter range, those features make it possible for structural modification and activity enhancement. With the beneficial biological effects shown in this report, MGF represents a novel pharmacophore that has the potential to be developed into a therapeutic improving metabolic syndrome.

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