A Metabolomics-driven Elucidation of the Anti-obesity Mechanisms of Xanthohumol\textsuperscript{a,\textdagger}

Jay S. Kirkwood\textsuperscript{b}, Lee Cole L. Legette\textsuperscript{b}, Cristobal L. Miranda\textsuperscript{b}, Yuan Jiang\textsuperscript{b}, and Jan F. Stevens\textsuperscript{b,\textdagger}

From the \textsuperscript{a}Linus Pauling Institute and the Department of Pharmaceutical Sciences and the \textsuperscript{b}Department of Statistics, Oregon State University, Corvallis, Oregon 97331


to whom correspondence should be addressed: Linus Pauling Institute, 307 Linus Pauling Science Center, Oregon State University, Corvallis, OR 97331. Tel.: 541-737-9534; Fax: 541-737-3999; E-mail: fred.stevens@oregonstate.edu.

Background: Xanthohumol may be beneficial for obesity-related conditions, but mechanisms are unknown.

Results: XN lowers ROS and dysfunctional lipid metabolism in animals, and XN uncouples respiration and induces oxidant defense systems in myocytes.

Conclusion: XN may ameliorate metabolic syndrome by these mechanisms.

Significance: Metabolomics helped reveal potential XN anti-obesity mechanisms.

Mild, mitochondrial uncoupling increases energy expenditure and can reduce the generation of reactive oxygen species (ROS). Activation of cellular, adaptive stress response pathways can result in an enhanced capacity to reduce oxidative damage. Together, these strategies target energy imbalance and oxidative stress, both underlying factors of obesity and related conditions such as type 2 diabetes. Here we describe a metabolomics-driven effort to uncover the anti-obesity mechanism(s) of xanthohumol (XN), a prenylated flavonoid from hops. Metabolomics analysis of fasting plasma from obese, Zucker rats treated with XN revealed decreases in products of dysfunctional fatty acid oxidation and ROS, prompting us to explore the effects of XN on muscle cell bioenergetics. At low micromolar concentrations, XN acutely increased uncoupled respiration in several different cell types, including myocytes. Tetrahydroxanthohumol also increased respiration, suggesting electrophilicity did not play a role. At higher concentrations, XN inhibited respiration in a ROS-dependent manner. In myocytes, time course metabolomics revealed acute activation of glutathione recycling and long term induction of glutathione synthesis as well as several other changes indicative of short term elevated cellular stress and a concerted adaptive response. Based on these findings, we hypothesize that XN may ameliorate metabolic syndrome, at least in part, through mitochondrial uncoupling and stress response induction. In addition, time course metabolomics appears to be an effective strategy for uncovering metabolic events that occur during a stress response.

Obesity results from an energy imbalance where intake exceeds expenditure. Given the increasing demand placed on health care systems by a rapidly growing obese population, there is great interest in developing a therapy to combat obesity and related disorders such as type 2 diabetes. To effectively reduce bodyweight and prevent obesity, therapeutic intervention must either increase energy expenditure or decrease energy intake or both. Mitochondria utilize macronutrients and downstream products to transport protons across the mitochondrial inner membrane and create an electrochemical potential, $\Delta \Psi$, which drives ATP synthesis. Uncoupling the dissipation of $\Delta \Psi$ from ATP synthesis allows for the unchecked catabolism of dietary macronutrients and formation of heat. In the early 20th century, the mitochondrial uncoupler 2,4-dinitrophenol was effectively used as an anti-obesity agent given to patients at doses ranging from 3 to 5 mg/kg daily (1). 2,4-Dinitrophenol was eventually not deemed safe for clinical use due to a narrow therapeutic window and serious risks from overdose, including death. However, given the effectiveness of general mitochondrial uncouplers and quickly rising rates of obesity worldwide, there is renewed interest in developing a safe alternative to 2,4-dinitrophenol.

Separate from expending energy, an increase in basal mitochondrial uncoupling reduces reactive oxygen species (ROS)\textsuperscript{2} generation and is associated with longevity (2–4). Mild, acute uncoupling increases the expression of the transcriptional coactivator, peroxisome proliferator-activated receptor-coactivator 1$\alpha$ (PGC-1$\alpha$), a dominant regulator of mitochondrial homeostasis in many tissues and a critical component of a cellular adaptive stress response (ASR) (5, 6). When a cell is challenged with an acute stress, it must quickly respond to counteract that stress to survive. The molecular adjustments made to cope with acute stress and the subsequent overcompensations together constitute an ASR. In the case of acute uncoupling, ATP levels drop but are normalized through increased expression of PGC1$\alpha$ (5). Low doses of toxic chemicals (e.g., 2,4-dinitrophenol) often activate ASR pathways but do not induce cell specific responses.

\textsuperscript{2}The abbreviations used are: ROS, reactive oxygen species; PGC-1$\alpha$, peroxisome proliferator-activated receptor-coactivator 1$\alpha$; ASR, adaptive stress response; XN, xanthohumol; DC, dicarboxylic; AC, acylcarnitine; HP, hydroperoxo; TXN, tetrahydroxanthohumol; BPNG, 8-prenylmaringenin; OCR, oxygen consumption rate; ECAR, extracellular acidification rate; TM, treatment medium; NAC, N-acetylcysteine; ETC, electron transport chain; UCP3, uncoupling protein 3; cySSG, cysteine glutathione disulfide; OP, ophthalmic acid; MnTMPyP, (manganese(III) tetrakis(1-methyl-4-pyridyl)porphyrin).
damage, allowing for a heightened ability to respond to subsequent stressors (7).

Mild ROS formation, reduced glutathione (GSH) depletion, and electrophilic stress are all triggers for induction of an ASR (8–10). In fact, there is recent evidence suggesting ROS and GSH depletion, under the right conditions, may actually prevent type 2 diabetes and increase longevity (11–14). Xanthohumol (XN) is a prenylated chalcone found in hops and beer and has an electrophilic group capable of inducing a stress response (10, 15). XN is perhaps best known for its cancer chemopreventive properties and its ability to trigger apoptosis through ROS (16, 17). Several reports are now starting to describe various anti-obesity properties of XN but with no well accepted mechanisms.

Our group recently reported that XN treatment decreased body weight gain and fasting plasma glucose in high fat fed Zuckerfa/fa rats, an established model of obesity and metabolic syndrome (18) (supplemental Fig. S1). In the present study, we sought to elucidate the mechanism(s) behind these changes using untargeted metabolomics. We analyzed the fasting plasma from the same Zucker rats treated with XN and found a global reduction in products of dysfunctional lipid metabolism, specifically dicarboxylic (DC) fatty acids and acylcarnitines (ACs). We also observed a decrease in hydroperoxy (HP) fatty acids, markers of oxidative stress. During fasting, DC fatty acids and ACs in plasma serve as reflections of incomplete fatty acid β-oxidation in skeletal muscle and are well characterized markers for assessing dysfunctional lipid metabolism (19, 20). These results prompted us to investigate the effects of XN on energy metabolism in muscle cells.

**EXPERIMENTAL PROCEDURES**

**Chemicals**—XN was isolated from female inflorescences of hops (Humulus lupulus) during previous work (21). Tetracydroxanthohumol (TXN) was also from previous work (22). 8-Prenylnaringenin (8PNG) was synthesized by prenylation of naringenin (23). MnTMPyP (manganese(III) tetrakis(1-methy1-4-pyridyl)porphyrin) and EUK 134 were from ENZO Life Sciences (Farmingdale, NY). Oligomycin A and N-acetylcysteine (NAC) were from Sigma. Metabolite standards were from TCI America (Portland, OR) and, when unavailable, Sigma.

**Zucker Rat Study**—Rat plasma utilized in this study was obtained from a previous investigation that examined the effects of chronic dosing of XN on metabolic syndrome (18). In brief, 4-week-old Zucker obese fa/fa male and female rats received daily oral doses of XN at 0, 1.86, 5.64, and 16.9 mg/kg body weight for 6 weeks. All rats were maintained on a high fat (60% kcal) AIN 93G diet for 3 weeks to induce severe obesity followed by a normal AIN-93G (15% kcal fat) diet for the last 3 weeks of the study. Plasma was collected at sacrifice via a carotid- puncture after fasting overnight. All procedures were approved by and in accordance with ethical standards of Oregon State University’s Institutional Animal Use and Care Committee (Protocol #3689).

**LC-MS/MS-based Metabolomics**—To extract plasma for metabolomics, plasma samples were thawed on ice. 80 μl of plasma was aliquoted into tubes, 240 μl of ice-cold 50:50 (v/v) MeOH:EtOH was added, and tubes were quickly vortexed 4 s. Samples were spun for 15 min at 13,000 × g at 4 °C, and the supernatant was analyzed by LC-MS/MS. To extract myocytes for time course metabolomics, media was aspirated, and cells were quickly washed once with warm Hanks’ balanced salt solution. After aspirating Hanks’ balanced salt solution, 400 μl of ice-cold 50:50 (v/v) MeOH:EtOH was quickly added to each well (6-well plate), then the plate was covered and placed in −80 °C freezer overnight. Cells were scraped then spun down for 15 min at 13,000 × g at 4 °C, and the supernatant was analyzed by LC-MS/MS. Metabolic profiling was performed as previously described, except column temperature was held at 50 °C (24). Importantly, the metabolite identification process was rigorous. Most metabolites were identified by mass, isotope distribution, MS/MS fragmentation pattern, and when standards were available, retention time. Further details can be found in the supplemental information.

**Cell Culture and Treatment**—Mouse C2C12 skeletal muscle cells and preadipocytes were first propagated in 75-cm² flasks using a culture medium consisting of DMEM (Invitrogen), 10% fetal bovine serum, 100 units/ml penicillin, 100 μg/ml streptomycin, and 1 mM pyruvate. Brown preadipocytes were a kind gift from Dr. Bruce Spiegelman (Harvard Medical School) and harvested as described in Uldry et al. (25). 3T3-L1 preadipocytes and C2C12 cells were from ATCC (ATCC, Manassas, VA). For metabolomics experiments, C2C12 myocytes were plated in 6-well plates at a density of 500,000 cells per well. 24 h later the medium was replaced with fetal bovine serum (FBS)-free XF medium (Seahorse Bioscience, North Billerica, MA) with 10 mM glucose (25 mM glucose for preadipocytes) and 1 mM pyruvate, and cells were allowed to incubate 1 h at 37 °C. Cells were then treated with XN or vehicle, 0.5% EtOH. For all cell based assays, XF medium was replaced with complete medium 4 h after XN addition to maintain cell viability for later time points. The final concentration of FBS after XN treatment was 0.1%. For cell viability experiments, the cells obtained from the culture flask after trypsinization were seeded in 96-well plates at a density of 10,000 cells per well using the same growth medium. After 24 h of incubation at 37 °C in a humidified atmosphere of 5% CO₂, the medium was removed, and the cells were rinsed once with Hanks’ balanced salt solution (Sigma). Freshly prepared XF assay medium containing 10 mM glucose and 1 mM pyruvate was then added to each well, 180 μl/well. After 1 h of incubation at 37 °C, 20 μl of culture media (9 parts XF assay medium and 1 part complete DMEM medium) containing various concentrations of XN were added to the wells. Ethanol was added to the control wells not to exceed a final concentration of 0.5%. The final concentrations of XN were 2, 5, 8, 25, and 70 μM, all in quadruplicate wells, in the 96-well plate.

**Respiration Measurements**—Oxygen consumption rate (OCR) and extracellular acidification rate (ECAR) measurements were performed with a Seahorse XF 24 Analyzer (Seahorse Bioscience). Cells were plated 24 h before measurements at a density of 20,000 cells per well for C2C12 myocytes and preadipocytes and 30,000 cells per well for HepG2 hepatocytes. The ECAR measurements are highly sensitive to FBS in growth medium, and it is suggested that little or none be present. XN is highly nonpolar, and previous experiments suggest FBS should be...
present in media to dissolve XN. To overcome this problem, XN was first dissolved in vehicle (ethanol), and the resulting solution was diluted 20-fold in treatment medium (TM) (9 parts XF media and 1 part 10% FBS media) that was added directly to cells. The TM contained 1% FBS, and after a further 10-fold dilution into the media in wells, only 0.1% FBS was present during OCR and ECAR measurements. This amount appeared to dissolve up to 70 μM XN at 37 °C and was well below the 1% limit recommended by Seahorse Bioscience. This treatment procedure was also used for oligomycin A, and accordingly, nearly 0.2% FBS was present in experiments using both. The three antioxidants were dissolved directly into serum-free XF medium before addition to cells. XN, TXN, 8PNG, oligomycin A, MnTMPyP, EUK 134, and NAC were prepared as fresh solutions the day of each experiment.

Statistical Analysis—Statistical analyses were performed using GraphPad Prism 4.0 and the software environment, R. In regard to the statistical analysis for the Zucker rat metabolomics study, the proportional odds model used to assess the variation of the metabolite intensities across the four levels was used to evaluate the significance of the metabolite variation across the four groups. The p values reported here arise from this analysis. Metabolites with a p < 0.01 from this test were considered metabolites of interest and are represented graphically with % change, XN 16.9 mg/kg versus control Zucker rats. Fisher’s method was used to combine individual p values and to test whether a group of metabolite intensities vary across different treatment groups. All acylcarnitines includes all carnitines represented here except carnitine and dehydroxy (DH) carnitine. Medium chain only includes only acylcarnitines 6:0, 8:0, 8:1, and 10:0. C, XN treatment decreased fasting plasma DC fatty acids and medium chain acylcarnitines, metabolites that result from dysfunctional lipid oxidation.

RESULTS

XN Reduces Products of Dysfunctional Lipid Metabolism and ROS—Several studies suggest that XN treatment could be beneficial for obesity, metabolic syndrome, and type 2 diabetes. Our group recently performed a 6-week treatment study with XN to obese Zucker rats and found attenuated body weight gain and decreased fasting plasma glucose in treated rats (supplemental Fig. S1) (18). To investigate the mechanism(s) of action responsible, we performed an untargeted metabolomics experiment on fasting plasma from the Zucker rats treated with XN at four different doses: 1) 0 mg/kg/day (vehicle control), 2) 1.86 mg/kg/day (low dose), 3) 5.64 mg/kg/day (medium dose), and 4) 16.9 mg/kg/day (high dose). XN was administered orally once per day. The animal doses correspond to doses of 0, 20, 60, and 180 mg/day for a human weighing 64 kg (27). Metabolites of interest are shown in Fig. 1A. XN decreased fasting plasma lev-
els of DC fatty acids, HP fatty acids, and one phosphatidylcholine (PC) (Fig. 1A). Amino acids were generally elevated in the high XN treatment group, but this change was specific to female rats (supplemental Fig. S2A).

DC fatty acids in the plasma and urine are markers of dysfunctional or substrate-overloaded mitochondria during fasting. With XN treatment, we report decreases in six different DC fatty acids varying in chain length and degrees of unsaturation (Fig. 1A). Five of the six were decreased >40% compared with controls (Fig. 1A). HP fatty acids are well characterized products of ROS and precursors to reactive lipid aldehydes. XN decreased the HP fatty acids 18:2 and 18:3 by 40 and 30%, respectively, suggesting lower levels of ROS production or an enhanced capacity to scavenge ROS in treated rats (Fig. 1A).

Although DC fatty acids are associated with dysfunctional mitochondrial β-oxidation, plasma ACs are often measured instead, as their detection is sensitive with mass spectrometry due to a constitutive positive charge and because they offer easy identification through a common, dominant, collision-induced fragment near m/z 85.028. We were able to target and identify 30 previously characterized ACs in our metabolomics data and, in agreement with the DC fatty acids, found that most were decreased with XN treatment (Fig. 1B). Although no single AC had a p < 0.01 from a Student’s t test, the comprehensiveness of the decrease could not be ignored. Fisher’s grouping analysis revealed a significant decrease in global (p = 0.0009) as well as medium chain ACs (p = 0.0091) (Fig. 1B). Targeting free fatty acids, precursors to ACs, in the metabolomics data revealed a global downward trend, albeit insignificant (supplemental Fig. S2B). A heat map, generated from correlations between metabolite levels across the four dose levels by computing the pairwise Pearson’s correlation, displayed a high correlation among ACs and among DC fatty acids highlighting the concerted role in the XN-induced OCR decrease, implicating superoxide or peroxynitrite as an antioxidant. In C–E, data are represented as the mean ± S.E., n = 4.

**XN Increases OCR and at High Concentrations Decreases OCR through ROS**—Electrons are transferred along the mitochondrial electron transport chain (ETC) where they ultimately split molecular oxygen (O₂) and water is formed. The energy gained from this process is used to transfer protons and generate ΔΨ. Mitochondria are responsible for 70–90% of total cellular O₂ consumption, and as such changes in cellular O₂ consumption largely reflect changes in mitochondrial O₂ consumption. At 5 µM, XN increased the OCR in C2C12 mouse skeletal muscle cells by roughly 30% compared with controls (Fig. 2A). XN (5 µM) also led to an increase in the ECAR, a measure of glycolytic ATP production (Fig. 2B). Together, these findings suggest a compensatory increase in energy generation in response to XN treatment.

Interestingly, at concentrations above 8 µM, XN inhibited OCR and ECAR almost completely (Fig. 2, C and D). When these cells were treated with XN at concentrations between 5 and 8 µM, inhibition of OCR was moderate, and ~50% inhibition was achieved with 6.5 µM (supplemental Fig. S3A). ECAR was still completely inhibited after 3 h at XN concentrations above 5 µM; however, the rate of ECAR decline was slower at lower concentrations, and 2 h post XN treatment, a noticeable dose-dependent inhibition with an IC₅₀ close to 6.5 µM could be observed (supplemental Fig. S3B). Although some studies suggest flavonoids inhibit electron transport by directly binding to ETC complexes, it has also been shown that XN promotes mitochondrial superoxide formation and, separately, that ROS can inhibit ETC activity (17, 28). To determine if ROS played a role in the XN-induced OCR decrease, myocytes were pretreated with antioxidants. The superoxide dismutase (SOD)/peroxynitrite dismutase mimetic MnTMPyP and the SOD/cat-alase mimetic EUK 134 both prevented the acute decrease in OCR caused by 70 µM XN and allowed for a substantial increase (Fig. 2E). However, the well characterized antioxidant NAC (a relatively poor scavenger of superoxide) did not prevent the decrease in OCR, implicating superoxide or peroxynitrite as

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**TABLE S1.** Because muscle tissue is the major consumer of fatty acids during fasting, we hypothesized the decrease in plasma ACs and DCs was a reflection of altered muscle metabolism. To test this, we treated muscle cells with XN and used bioenergetic and metabolomic approaches to uncover its effects.

**FIGURE 2.** XN has a hormetic effect on respiration and glycolytic energy production. A, 5 µM XN acutely increases OCR in C2C12 mouse myocytes by 25% (p = 0.0037, Student’s two-tailed t test, n = 10). B, 5 µM XN decreased the ECAR by 20% (p = 0.0041, 3 h post treatment, Student’s two-tailed t test, n = 10). C, shown is the dose-dependent effect of XN on myocyte OCR. D, shown is the dose-dependent effect of XN on myocyte ECAR 2 h post treatment. E, shown is the effect of antioxidant pretreatment on the 70 µM XN induced decrease in OCR; An, antioxidant. In C–E, data are represented as the mean ± S.E., n = 4.
possible culprits (Fig. 2E). Neither EUK 134 (50 μM) nor MnTMPyP (50 μM) affected OCR, but a relatively low (compared with other studies) concentration of NAC (0.5 mM) was used because higher concentrations greatly decreased respiration (supplemental Fig. S4). These results suggest XN inhibits the ETC and OCR through the production of ROS.

**XN Increases Uncoupled Respiration**—The majority of O₂ consumed inside cells is coupled to ATP synthesis. To test whether the increase in OCR due to XN was coupled to ATP production, we pharmacologically inhibited ATP synthase with oligomycin A before XN treatment. In the presence of oligomycin A, XN increased OCR 2-fold, indicative of mitochondrial uncoupling (Fig. 3A). This effect was even more pronounced at 70 μM XN but only with antioxidant pretreatment (Fig. 3B).

Mitochondria are present in almost every cell type and provide a ubiquitous target for energy expenditure. As such, non-selective mitochondrial uncouplers increase respiration in most cell types limited largely by their ability to reach their target. We treated HepG2 hepatoma cells as well as brown and white preadipocytes with XN to assess the effect on OCR in different cell lines. XN increased OCR in HepG2 cells but only when pretreated with an antioxidant, perhaps due to a heightened sensitivity to ROS in this cell line compared with C2C12 myocytes (Fig. 3, C and D). But similar to the myocytes, the increase in respiration in HepG2 cells was not coupled to ATP synthesis (Fig. 3E). XN also increased uncoupled respiration in white and brown preadipocytes (Fig. 4A). And again, at higher concentrations, XN inhibited respiration through ROS (Fig. 4, B and C).

**XN Electrophilicity Is Not Necessary for Effects on Respiration**—Uncoupling protein 3 (UCP3) and adenine nucleotide translocase are both mitochondrial proteins that contribute to a basal level of proton leak and mitochondrial uncoupling in muscle (29). Both possess exposed thiol groups that can affect enzyme activity upon modification (1). UCP3 activity is in part regulated by glutathionylation and adenine nucleotide translocase by thiol oxidation (30, 31). Furthermore, the electrophilic lipid peroxidation product, 4-hydroxy-2(3C)-nonenal, can uncouple respiration, likely through UCPs and adenine nucleotide translocase (32). Because XN also has an electrophilic group capable of thiol alkylation, we hypothesized that XN may act in a manner similar to 4-hydroxy-2(3C)-nonenal. To test this idea, we utilized the hydrogenated product of XN, TXN, which lacks the electrophilic α,β-unsaturated ketone functionality of XN (Fig. 5A). Treatment of myocytes with TXN resulted in an increase in OCR at low concentrations and a decrease at higher concentrations, similar to the hormetic dose response observed with XN (Fig. 5B). However, TXN was more effective at increasing respiration compared with XN, resulting in a similar increase in OCR but at less than half the concentration. 8PNG is an estrogenic in vivo metabolite of XN that had little effect on myocyte OCR, even at 12 μM (Fig. 5C).

**Time Course Metabolomics Reveals a Catabolic Phenotype and Induction of an ASR**—XN (5 μM) uncoupled respiration and activated glycolysis in myocytes for several hours, indicating a highly catabolic state inside the cell (Fig. 2, A and B). To capture the associated, acute metabolic changes as well as long term changes in metabolism, we performed untargeted, time course metabolomics on myocytes treated with XN (5 μM). To provide an accurate correlation between the myocyte metabolome and the previous respiration measurements, experimental conditions for the two separate assays were kept similar.
Serum-free medium is required for the OCR/ECAR assays and was, therefore, also used for the metabolomics experiments. However, serum-free medium was replaced with 10% FBS medium at 4 h post-treatment to maintain cell viability for later time points (Fig. 6A). Cells were extracted at eight different time points after XN addition, from 10 min to 48 h (Fig. 6A). Results were not normalized because cell viability and cellular protein content were unaffected by concentrations less than 25 μM XN (supplemental Fig. S5, A and B). Significantly differing metabolites (Student’s t test, control versus 5 μM XN, >2-fold change, \( p < 0.01 \), for any time point) are displayed at their most significant time point (time point with greatest -fold change with \( p < 0.01 \)) in Fig. 6B.

The largest change observed in XN-treated cells was a 10.5-fold increase in GSH (Fig. 6B). GSH represents the main endogenous antioxidant inside cells, and when ROS increases, GSH is consumed, and oxidized glutathione (GSSG) is produced (33). GSSG is recycled enzymatically back to GSH in a NADPH-dependent manner to maintain the GSH/GSSG ratio. After XN addition, we observed an acute increase in GSH as well as a decrease in GSSG, suggesting acute activation of GSSG recycling systems (Fig. 7A). At 16 h, GSH and GSSG were elevated 10.5- and 1.7-fold respectively, indicating up-regulated GSH synthesis separate from recycling (Fig. 7B). The GSH/GSSG ratio was elevated in XN-treated myocytes during the entire experiment (Fig. 7C). In contrast, cysteine glutathione disulfide (cySSG) was decreased during the entire experiment (Fig. 7C). cySSG is a product of ROS, cysteine, and GSH but may also serve as a GSH reservoir in times of need (34, 35). We observed a highly significant, inverse correlation between cySSG and GSH, but only in XN-treated cells, supporting this notion (Fig. 7D).

Methionine is highly sensitive to oxidation by ROS. Its mild oxidation product, methionine sulfoxide (Met-SO), is enzymatically recycled back to methionine by methionine sulfoxide reductase (Msr) in a thioredoxin (Trx)-dependent manner. Therefore, Met-SO may serve as a sensitive marker for acute induction of ROS and their effective removal (Fig. 7E) (36). XN induced a transient increase in Met-SO with maximal levels at the 4-h time point (Fig. 7F and G). The increase was less apparent when normalized to free methionine but still significant (Student’s t test, \( p < 0.001 \)) (Fig. 7F).
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The electrophilicity of XN and the observed changes in glutathione metabolism prompted us to search for a XN-glutathione adduct (XN-SG). XN-SG was detected and identified by thiosulfate metabolism prompted us to search for a XN-glutathione adduct (XN-SG). XN-SG was detected and identified by mass spectrometry. The mass, isotope distribution, and fragmentation pattern of XN-SG were characterized and compared to that of a standard glutathione adduct (GSH-SG). The time course profile of XN-SG was similar to that of GSH-SG, with a peak at 4 h after XN treatment (Fig. 8A). Six peptides (tentatively identified by MS as tripeptides and dipeptides) in total were uncovered as significantly differing metabolites (Fig. 8, A and B). Interestingly, all six increased the first 4 h after XN treatment but were sharply decreased at 24 h (Fig. 8, A and B).

We observed an acute decrease in the active B vitamin precursors thiamine and pyridoxine after XN treatment (Fig. 8C). Thiamine and pyridoxine levels were positively correlated in treated and untreated samples (Fig. 8D). For most time points, both of these B vitamins were lower in XN-treated cells, and accordingly, the treated samples occupy the lower left portion of the plot (Fig. 8D). Folic acid and nicotinamide were also lower in XN-treated myocytes, whereas pantothenic acid was unchanged (Fig. 8E).

It may also be worth mentioning that adenosine phosphates were altered by XN. ADP was elevated 4.7-fold 16 h after XN treatment (Fig. 8F). AMP levels were mildly altered (Fig. 8F).

The remaining findings, depicted in Fig. 9, are the result of a targeted data mining effort specifically focusing on metabolites central to energy metabolism. Metabolites targeted include those from glycolysis, the tricarboxylic acid (TCA) cycle, phosphocreatine metabolism, amino acids, and lipids (supplemental Fig. S7). The parameters used to identify individual metabolites can be found in supplemental Table S2. Overall, these changes appear to reflect a state of acute energy stress and hint toward induction of protein degradation (Fig. 9; supplemental Fig. S7). These changes and their implications are discussed in detail below.

**DISCUSSION**

The most commonly used plasma markers of dysfunctional \( \beta \)-oxidation in type 2 diabetes are ACs, thought to be byproducts of mitochondrial dysfunction and fatty acid overload in skeletal muscle (19, 38). During fasting, fatty acids are released into the bloodstream for uptake and mitochondrial \( \beta \)-oxidation by peripheral tissues (39). Skeletal muscle constitutes about 40% of total body mass and is the major consumer of fatty acids during fasting (40). The presence of dysfunctional or too few mitochondria in skeletal muscle will result in high rates of incomplete fatty acid oxidation and elevated levels of ACs in plasma (19, 41). Decreasing levels of these species will either alleviate mitochondrial dysfunction or their reduction serves as a marker of prior alleviation of such dysfunction. We found that XN-treated Zucker rats had globally decreased levels in plasma ACs, but individual ACs failed to reach statistical significance (Fig. 8C). Thiamine and pyridoxine levels were positively correlated in treated and untreated samples (Fig. 8D). Folic acid and nicotinamide were also lower in XN-treated myocytes, whereas pantothenic acid was unchanged (Fig. 8E).

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The most significant change in our metabolomics data, however, appears to be the decrease in DC fatty acids, also a marker of dysfunctional lipid metabolism although typically measured in the urine for diagnosis of inherited \( \beta \)-oxidation disorders (42) (Fig. 1A). DC fatty acids are endogenous products of cellular fatty acid metabolism, first \( \omega \)-hydroxylated by endoplasmic
reticulum-associated enzymes then further oxidized in the peroxisome (43). DC fatty acids are normally present at relatively low levels, but their production increases when fatty acid supply is excessive or mitochondrial β-oxidation is dysfunctional (44). DC fatty acids are able to undergo β-oxidation in the peroxisome or mitochondria or they can be excreted in the urine due increased hydrophilicity (43). The decrease in DC fatty acids and other lipid species with XN treatment strongly hints toward amelioration of mitochondrial dysfunction in skeletal muscle (Fig. 1C). Also, given the small degree of intergroup variation observed in DC fatty acids relative to ACs, fasting levels of DC fatty acids may serve as a more consistent indicator for mitochondrial dysfunction, especially when group size is small.

Hydroxylated chalcones have previously been implicated as mitochondrial uncouplers (45). Prenylated derivatives such as XN may be more effective uncouplers as they are highly non-polar and easily diffuse across lipid bilayers. This notion taken together with our metabolomics data prompted us to investigate the effects of XN on respiration and energy metabolism in muscle cells. We found that XN acutely increased uncoupled respiration in muscle cells, and at 5 µM this effect was significant (Figs. 2A and 3A). The effective concentration of XN in cell culture (2–5 µM) was similar to levels observed in Zucker rats (1–3 µM in liver, <1 µM in plasma) (18, 46). Interestingly, XN at 8 µM and above inhibited respiration in myocytes (Fig. 2, C and D). Pharmacological intervention with antioxidants MnTMPyP and EUK 134, but not NAC, completely prevented the decrease in OCR induced by 70 µM XN, hinting toward superoxide- or peroxynitrite-mediated inhibition of respiration (Fig. 2E). XN also increased uncoupled respiration in HepG2 hepatoma cells, but at higher concentrations, and only with antioxidant (MnTMPyP) pretreatment (Fig. 3, D and E). XN had a similar effect in white and brown preadipocytes as in myocytes; low levels increased uncoupled respiration, and high levels inhibited respiration in a ROS-dependent manner (Fig. 4). It was previously hypothesized that XN inhibits respiration in cancer cell lines by directly binding the Fe centers of the ETC (17). Our results suggest the inhibitory effects of XN on respiration may simply stem from ROS production.

To test if the electrophilicity of XN played a role in the OCR increase or decrease, we treated myocytes with TXN, a hydrogenation product of XN that is not electrophilic (Fig. 5A). TXN had a similar dose-response curve to that of XN but was equally effective at increasing OCR at less than half the concentration (Fig. 5B). TXN had a similar dose-response curve to that of XN but was equally effective at increasing OCR at less than half the concentration (Fig. 5B). This increased efficacy likely results from a lack in protein thiol interactions due to the absence of electrophilicity. From these results we conclude that the α, β-unsaturated ketone functionality of XN is not required for the acute effects on respiration. 8PNG is a potent phytosterogen and an in vivo metabolite of XN. However, 8PNG failed to have any effect on OCR in myocytes (Fig. 5C).

Post translational signaling events such as protein phosphorylation, glutathionylation, and thiol modification aid in the acute regulation of metabolism and respiration and help facilitate the stress response (47–50). To monitor all the modifications to proteins involved in energy metabolism would require an amazing effort. Metabolomics allows for a relatively easy correlation to cellular phenotype due to the lack of modifications, such as those made to proteins after translation (51). In
muscle cells, XN (5 μM) acutely uncoupled respiration and increased ECAR, reflecting an elevated state of energy production. To uncover the associated changes in metabolism, we performed a time course metabolomics experiment on myocytes treated with XN (Fig. 6A). Unexpectedly, glutathione recycling was immediately increased, and synthesis was greatly elevated 16 h after XN exposure, when GSH was increased 10.5-fold (Fig. 7, A and B). The oxidation product of GSH, cySSG, has previously been hypothesized to act as a reservoir for GSH (35). We observed a highly significant (p < 0.0001) correlation between cySSG and GSH but only in XN-treated cells (Fig. 7C). These results support the role of cySSG as a GSH reservoir during times of stress. Activation of the ASR requires Nrf2, which is critical for glutathione recycling and targeted degradation of damaged proteins (52). Because XN can prevent inactivation of Nrf2 via Keap1 adduction, these changes may be a reflection of increased Nrf2 activity. We were able to detect the XN-SG adduct, a marker of electrophilic stress (Fig. 7D).

ROS can also modify the sensitive Keap1 thiols allowing for Nrf2 release and activation. Our respiration experiments indicate that high concentrations of XN inhibit respiration through ROS. Metabolomics revealed that XN (5 μM) transiently increased Met-SO, a product of methionine and ROS (Fig. 7E). Met-SO was elevated 2-fold at 4 h but normalized by 10 h (Fig. 7F and G). To try and account for changes due to protein degradation, we normalized Met-SO to free methionine levels and found a slightly blunted but still significant (p < 0.0001) increase at 4 h (Fig. 7F). These findings, however, do not

FIGURE 8. XN affects protein degradation, cofactor, and energy metabolism. A, shown is the time course response of three tripeptides after XN treatment. All three tripeptides were significantly increased (p < 0.05) 4 h post XN treatment, and all were decreased (p < 0.01) at 24 h. B, shown is the time course response of three dipeptides after XN treatment. All three dipeptides were significantly decreased (p < 0.01) 24 h post XN treatment. C, shown is the time course response of pyridoxine and thiamine after XN treatment. D, shown is the correlation between thiamine and pyridoxine levels in XN-treated and control myocytes. All time points are included. E, shown is the time course response of folic acid, nicotinamide, and pantothenic acid after XN treatment. F, shown is the time course response of AMP and ADP after XN treatment. *, p < 0.05; **, p < 0.01; ***, p < 0.001, from Student’s two-tailed t test.
account for selective degradation of proteins with high levels of Met-SO.

Ophthalmic acid is a GSH analog lacking the thiol and is a potential marker of oxidative stress (37). Although OP and its analog norophthalmic acid were both found to be decreased long term when GSH was greatly elevated, their overall time course profiles were strikingly similar to those of several uncharacterized peptides, increasing roughly 2-fold over the first 4 h then dropping 3–4-fold below control levels 24 h post XN treatment (Fig. 8, A and B). In the context of this experiment, we believe these peptides are products of incomplete protein degradation, potentially both proteasomal and lysosomal. In general, peptides were initially increased (presumably to support the enhanced respiration) followed by an even greater decrease (presumably due to up-regulated protein synthesis and organelle biogenesis) (Fig. 8, A and B). In addition, given the consistency in the profiles of these peptides over time, we propose that their monitoring by LC-MS/MS may provide a small molecule-based method for the determination and tracking of cellular autophagy. Because the proteases in the cytosol and lysosomes have different amino acid specificities, we hypothesize differential di- and tripeptides may arise from the two processes. This idea is being explored.

ADP was increased ~4-fold 16 h post treatment (Fig. 8F). As an adaptation to exercise, the mitochondrial affinity for cytosolic ADP in muscle decreases as the myofibers become more oxidative (53). It is tempting to speculate that the long term increase we observed in ADP may be related to this switch.

Also of interest, we observed a decrease in several precursors to active B vitamins, cofactors necessary for the catabolism of macronutrients. Thiamine and pyridoxine are B vitamin precursors to the biologically active cofactors, thiamine pyrophosphate and pyridoxal phosphate. Thiamine pyrophosphate and pyridoxal phosphate are cofactors for many enzymatic reactions and are required for the degradation of fatty acids, sugars, and amino acids. We observed an acute decrease in these precursors with XN treatment, but their levels were correlated regardless of treatment, suggesting constitutive, coordinate regulation (Fig. 8D). Similar to the other B vitamins, nicotin-
amide and folic acid were lower in XN-treated myocytes over the first 4 h (Fig. 8E). We hypothesize that these changes are a reflection of up-regulated, active B vitamin production for use as cofactors in catabolic reactions.

An acute increase in OCR and ECAR, indicating a catabolic state within the cell, was accompanied by a variety of alterations in energy metabolism. Creatine kinase catalyzes the transfer of a high energy phosphate from phosphocreatine to ADP, resulting in ATP and creatine. The maximal rate of ATP synthesis through the creatine kinase system is much higher than oxidative phosphorylation, and as such, phosphocreatine serves as an energy reservoir that supports acute energy depletion (54). After XN treatment, time course metabolomics revealed activation of the acute compensatory creatine kinase system, indicated by a decrease in phosphocreatine at 10 min. Creatine did not increase until 4 h post treatment. However, the spontaneous intramolecular condensation product of creatine, creatinine, increased immediately ($p > 0.05$). Lactate production from pyruvate also produces ATP and is another mechanism for muscle to quickly recover from energy depletion. Although lactate and pyruvate were both decreased for the first 4 h ($p > 0.05$), we observed a sustained increase in ECAR, a measure of cellular lactate export and glycolytic activity (Fig. 2B). Furthermore, intracellular hexose (our method cannot distinguish glucose from other hexoses) was decreased short term, perhaps due to enhanced glucose utilization.

Under conditions of energy depletion, cells maintain energy status by selectively degrading macronutrient complexes including proteins and organelle membranes, a process known as autophagy. In XN-treated myocytes, metabolomics revealed a transient increase in several uncharacterized di- and tripeptides as well as the majority of amino acids, indicative of protein degradation and amino acid mobilization for entry into the TCA cycle. Free glutamine is present in muscle at high concentrations and is a major amino acid feeding the TCA cycle (55, 56). Glutamine is first converted to glutamate, which can then produce the TCA cycle intermediate oxoglutarate. Interestingly, glutamine was the only amino acid that significantly decreased in the first 4 h, perhaps due to a higher rate of utilization than production (supplemental Fig. S7).

To undergo $\beta$-oxidation, fatty acyl CoA species are converted to ACs for transport into the mitochondria. Plasma ACs increase during exercise due to an elevation in fatty acid supported respiration in muscle (57). Again in agreement with a state of acutely increased respiration, we observed modest increases in both acetylcarnitine and long chain ACs in the first 4 h post treatment. Possibly due to increased demand and synthesis, carnitine was elevated after 4 h. Perhaps most importantly, these results highlight the dynamic nature of the cellular stress response and suggest metabolomics may be an effective strategy for studying this response.

Together, our cell culture experiments suggest XN has three general, acute effects: mitochondrial uncoupling, ROS generation, and electrophilic adduction (Scheme 1). Although the ROS production and electrophilic adduction are previously documented, the uncoupling activity of XN has not been. Separately, these mechanisms are capable of activating an ASR, and their simultaneous occurrence, as observed with XN treatment, may ensure this induction. For example, Nrf2 is activated by both ROS and electrophilic adduction (Scheme 1).

| ACUTE EFFECTS | DOWNSTREAM EFFECTS |
|---------------|---------------------|
| **XN** | **XN** |
| Electrophilic Adduction | ROS scavenging capacity |
| ROS Generation | Catabolism |
| Mitochondrial Uncoupling | Autophagy |
| Energy expenditure | Energy expenditure |
| Mitochondrial biogenesis | Oxidative capacity |
| PGC1α | |

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mitophagy, and increased mitochondrial biogenesis and oxidative capacity. We propose that the beneficial effects observed in XN-treated Zucker rats stem at least in part from these three general mechanisms and subsequent activation of adaptive cellular processes (Scheme 1).

Although XN treatment reduced Zucker rat body weight gain in a previous study by our group, it is important to note this change was only significant in males (supplemental Fig. S1). Because protonophoric mitochondrial uncouplers act directly, this result does not appear to fall in line with this proposed action of XN. However, it is a well-documented phenomenon that females have an enhanced response to a high fat feeding by increasing UCP expression in muscle and brown adipose tissue (58, 59). Importantly, enhanced UCP expression could lower mitochondrial membrane potential and decrease the effectiveness of weakly acidic mitochondrial uncouplers, where activity is driven in part by membrane potential and pH (60–62). We propose that enhanced UCP expression in the female Zucker rats may have blunted the effects of XN on body weight. However, the decrease in plasma markers of dysfunctional lipid metabolism described in this study were not found to be sex-specific, suggesting these changes may not be related to the protonophoric properties of XN. Instead, we hypothesize they are a reflection of an adaptive cellular response to electrophilic and oxidative stress induced by XN (Scheme 1).

Although XN clearly uncouples respiration in cell culture, our experiments do not rule out other mechanisms that may be responsible for the effects of XN in Zucker rats. Alternatively, we propose that XN may exert its beneficial effects through increased bile acid production. Plasma bile acids induce energy expenditure by activating the G-protein-coupled receptor, TGR5, which promotes intracellular thyroid hormone activation and, subsequently, UCP expression in both muscle and brown adipose tissue (63, 64). Although no reports describe a direct increase in plasma bile acids with XN treatment, at least two hints strongly toward it. In 2005, Nozawa (65) reported a 7-fold induction in the mRNA of CYP7A1 in the livers of mice fed XN. CYP7A1 hydroxylates cholesterol, and this is the rate-limiting step in bile acid synthesis. In 2012, Hirata et al. (66) saw an increase of bile acids in the feces of XN-treated mice. Neither study reported bile acid measurements in plasma. However, the large diurnal and interindividual variation in plasma bile acid concentrations makes comparative measurements difficult (67). In agreement with this, targeted data mining of bile acids revealed great variation between XN-treated and control Zucker rats, and no significant changes were observed (data not shown).

CONCLUSIONS

In vivo, metabolomics revealed that XN treatment markedly and globally reduced plasma markers of ROS and dysfunctional lipid oxidation in the muscle tissue of Zucker rats. In vitro we show that XN uncouples respiration in several cell types, and metabolomics revealed an adaptive stress response in myocytes. The degree to which these mechanisms occur and contribute to the anti-obesity effects of XN in vivo is currently under investigation.

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