Increasing NAD Synthesis in Muscle via Nicotinamide Phosphoribosyltransferase Is Not Sufficient to Promote Oxidative Metabolism*

David W. Frederick†, James G. Davis‡, Antonio Dávila, Jr.¶, Beamon Agarwal‡, Shaday Michan§, Michelle A. Puchowicz**, Eiko Nakamaru-Ogiso**, and Joseph A. Baur††

From the ‡Department of Physiology, Institute for Diabetes, Obesity, and Metabolism and †Department of Biochemistry and Biophysics, Perelman School of Medicine, University of Pennsylvania, Philadelphia, Pennsylvania 19104, §Instituto Nacional de Geriatría, México, Distrito Federal 10200, México, and **Department of Nutrition, Mouse Metabolic Phenotyping Center, Case Western Reserve University, Cleveland, Ohio 44106

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† To whom correspondence should be addressed: Dept. of Physiology, Inst. for Diabetes, Obesity, and Metabolism, Perelman School of Medicine, University of Pennsylvania, 3400 Civic Center Blvd., SCTR 12-114, Philadelphia, PA 19066. Tel.: 215-746-4585; Fax: 215-898-5408; E-mail: baur@mail.med.upenn.edu.

The NAD biosynthetic precursors nicotinamide mononucleotide and nicotinamide riboside are reported to confer resistance to metabolic defects induced by high fat feeding in part by promoting oxidative metabolism in skeletal muscle. Similar effects are obtained by germ line deletion of major NAD-consuming enzymes, suggesting that the bioavailability of NAD is limiting for maximal oxidative capacity. However, because of their systemic nature, the degree to which these interventions exert cell- or tissue-autonomous effects is unclear. Here, we report a tissue-specific approach to increase NAD biosynthesis only in muscle by overexpressing nicotinamide phosphoribosyltransferase, the rate-limiting enzyme in the salvage pathway that converts nicotinamide to NAD (mNAMPT mice). These mice display a ~50% increase in skeletal muscle NAD levels, comparable with the effects of dietary NAD precursors, exercise regimens, or loss of poly(ADP-ribose) polymerases yet surprisingly do not exhibit changes in muscle mitochondrial biogenesis or mitochondrial function and are equally susceptible to the metabolic consequences of high fat feeding. We further report that chronic elevation of muscle NAD in vivo does not perturb the NAD/NADH redox ratio. These studies reveal for the first time the metabolic effects of tissue-specific increases in NAD synthesis and suggest that critical sites of action for suplemental NAD precursors reside outside of the heart and skeletal muscle.

Nicotinamide adenine dinucleotide (NAD) is a ubiquitous pyridine nucleotide that functions as an essential reduction-oxidation (redox) cofactor in cellular metabolism. Recent years have seen growing interest in the expanded roles for NAD as a co-substrate in a myriad of signaling contexts, spurred in part by the discoveries that in yeast enhancing flux through the NAD salvage pathway is necessary and sufficient for lifespan extension by calorie restriction (1, 2) and that in higher eukaryotes NAD levels fluctuate in response to circadian cues, exercise, and nutritional status (3–5). In mammalian cells, NAD is continuously consumed as a substrate in adenosine diphosphate (ADP)-riboseylation, cyclization, and deacylation reactions that influence a host of physiological processes (6). These functions highlight the necessity for NAD biosynthetic pathways in the maintenance of cellular homeostasis and suggest that modulation of the NAD pool might have therapeutic applications.

Although the de novo synthesis of NAD from dietary tryptophan, nicotinic acid, or nicotinamide-containing nucleotides can contribute to the intracellular NAD pool in mammals, the constitutive degradation of NAD to nicotinamide (NAM) by endogenous enzymes renders a functional NAD salvage pathway necessary for life (7). The rate-limiting enzyme in the NAD salvage pathway, nicotinamide phosphoribosyltransferase (NAMPT/NamPTase/PBEF/Visfatin), catalyzes the condensation of NAM and phosphoribosyl pyrophosphate to nicotinamide mononucleotide (NMN), which is subsequently converted to NAD by NMN adenyltransferases (NMNAT1–3) (8).

NAD salvage is at least partly regulated at the level of NAMPT protein expression, which increases in response to various stresses and disease states, including inflammation,
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hypoxia, and oncogenesis. NAMPT overexpression is sufficient to elevate NAD concentration and confer cytoprotective effects upon cells in culture (8–10) and in vivo (11). In skeletal muscle, NAMPT protein levels have been observed to more than double in response to chronic exercise (12, 13) or calorie restriction (14, 15). Importantly, administration of the specific NAMPT inhibitor FK866 to calorie-restricted rats can partially or fully revert several beneficial metabolic phenotypes to the ad libitum conditions (15). Although NAMPT maintains a critical co-substrate of the sirtuin family of deacylases and ADP-ribosyltransferases (16), the range of mechanisms through which it can influence mammalian physiology remains largely unresolved.

The observations that global increases in NAD availability appear beneficial to mammalian metabolic health, especially in the context of nutrient excess, do not reveal the particular cells or tissues responsible for the effects. Whole body deletion of major NAD consumers, including CD38 (17), PARP-1 (18), and PARP-2 (19), provides protection against diet-induced obesity and metabolic disease, although it is difficult to determine whether these phenotypes derive from the increased availability of NAD versus the loss of signals that would otherwise arise from the deleted enzymes. Studies targeting NAD synthesis via the direct provision of the NAMPT product, NMN, as well as its unphosphorylated relative, nicotinamide riboside (NR), have bolstered the case that higher intratissue NAD levels improve oxidative metabolism and ameliorate deleterious consequences of a high fat diet (20, 21). Such NAD-boosting strategies have been associated with enhanced mitochondrial biogenesis and function in tissues where elevated NAD was observed, including liver (21), brown adipose (20), and skeletal muscle (17). These findings coincide with enhancements in energy expenditure, thermogenesis, and endurance (19, 20) as well as correction of respiratory and ultrastructural defects in mouse models of mitochondrial disorders (22, 23). However, NAD-modulating interventions involving either germ line genetic mutations or transient, systemically administered treatments cannot conclusively resolve the primary site of action for NAD or the contributions of individual tissues to the whole-animal metabolic phenotype. Thus, a conditional transgenic approach is needed to specifically and chronically modulate NAD homeostasis in metabolically significant tissues.

In this study, we investigated the degree to which increasing NAD concentration exerts tissue-autonomous effects on oxidative metabolism in otherwise healthy muscle. By specifically overexpressing NAMPT in muscle, we provide evidence that a modest increase in the steady-state level of NAD is insufficient to improve muscle oxidative function or ameliorate consequences of high fat diet feeding and does not alter the NAD/NADH ratio. The lack of phenotype in our transgenic model suggests that NAD-boosting treatments may benefit muscle via signals that originate from outside the tissue.

**EXPERIMENTAL PROCEDURES**

**Generation of Muscle-specific Nampt Transgenic Mice**—A Cre-inducible Nampt construct was generated by placing a floxed transcriptional stop element between a CAGGS promoter and the Nampt cDNA. This construct was then targeted to the Col1A1 locus as described previously (24). Nampt<sup>fl/fl</sup>/WT mice were generated by crossing Nampt<sup>fl/fl</sup>/WT or Nampt<sup>fl/fl</sup>/H11001 mice to muscle creatine kinase (MCK)-Cre transgenic mice on the C57BL6/J background obtained from The Jackson Laboratory. Nampt<sup>+/+</sup> mice were subsequently generated by crossing Nampt<sup>+/+</sup>/WT to Nampt<sup>fl/fl</sup>/WT or Nampt<sup>fl/fl</sup>/H11001 mice. Littersmates lacking either MCK-Cre or the floxed allele were used as controls. Mice were studied between the ages of 3 and 9 months as indicated. The NAM supplementation experiment in Fig. 11 was performed in 15-month-old animals.

**Animal Housing and Diets**—All mice were housed according to approved Institutional Animal Care and Use Committee protocols and sacrificed after an overnight fast. Animals were kept in a specific pathogen-free barrier facility with controlled temperature and humidity and a 12-h/12-h light/dark cycle with lights on at 7 a.m. Mice were fed standard chow diet containing 120 ppm niacin (Rodent Diet 5010, LabDiet) unless otherwise noted. Obesity was induced with high fat diet (HFD) (D12492, Research Diets) containing 60% calories from fat. For NAM and NR supplementation experiments, compounds were dissolved weekly in the drinking water at 2.67 mg/ml in light-protected bottles.

**Immunoblotting**—Tissues were lysed with a motorized homogenizer in lysis buffer (25 mM Tris-HCl, pH 7.9, 100 mM KCl, 5 mM MgCl<sub>2</sub>, 1% (v/v) Nonidet P-40, 10% (v/v) glycerol) supplemented with Complete protease inhibitor mixture (Roche Applied Science). SDS-PAGE was performed on 4–15% Tris-HCl gels (Bio-Rad) using 20 μg of cleared lysate boiled for 5 min in 1× Laemmli buffer. Gels were transferred to PVDF membranes (Millipore) and blocked with 5% nonfat milk. Primary antibodies were diluted 1:1000 in TBS-Tween 20 according to the manufacturer’s instructions. PBEF (A300-372A) was from Bethyl Laboratories, Mitoprofile Total OXPHOS antibody mixture (ab110413) was from Abcam, and β-actin-HRP (ab49900) was from Abcam. Antibody binding was detected using chemiluminescent horseradish peroxidase substrate (PerkinElmer Life Sciences) and quantified using NIH Imagej software.

**RNA and DNA Extraction**—RNA was isolated from ~25 mg of muscle tissue using 1 ml of TRIzol (Qiagen) reagent according to the manufacturer’s instructions. For total DNA isolation, ~25 mg of tissue was incubated in digestion buffer (100 mM Tris-HCl, pH 8.5, 200 mM NaCl, 5 mM EDTA, 0.2% (w/v) SDS, 1 mM β-mercaptoethanol) supplemented with 0.3 mg/ml protease K (Sigma) overnight at 55 °C followed by an additional 0.3 mg/ml protease K for 1 h at 55 °C. Nucleic acids were precipitated by addition of 1.25 mM NaCl and ethanol, washed in 70% (v/v) ethanol, and resuspended in Tris-EDTA buffer supplemented with 80 μg/ml RNase A (Roche Applied Science).

**Quantitative Real Time PCR and Recombination Analysis**—Total RNA (1 μg) was reverse transcribed using a High Capacity cDNA Reverse Transcription kit (Applied Biosystems). Quantitative real time PCR for gene expression was performed on an Applied Biosystems 7900HT system using 15 ng of cDNA/reaction and a standard curve method relative to reference gene 36B4. One control sample was excluded as an outlier according to Grubb’s test with α = 0.05 due to low reference gene expression. For LoxP recombination analysis, 15 ng of total DNA was analyzed by a custom TaqMan Copy Number
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Assay (Invitrogen) normalized to reference gene Trif with custom primers flanking the 3’ LoxP site of the transgene. For determination of mitochondrial DNA (mtDNA)/nDNA content, 15 ng of total DNA was analyzed with primers specific to mitochondrial gene ND1 (forward, 5-TAA CGC GAA TGG GCC GGC TG-3; reverse, 5-TAA CGC GAA TGG GCC GGC TG-3) and nuclear gene TBP (forward, 5-CCT GTA CCC TTC ACC AAT-3; reverse, 5-GAA GCT GCG GTA CAA TTC CAG-3).

Glucose Tolerance Test—Mice were injected intraperitoneally with d-glucose solution at a dose of 1.5 g/kg of body weight after an overnight fast. Injected mice were allowed free locomotion and access to water. Tail vein blood glucose was measured with a handheld glucometer (Abbott) for the next 2 h.

Citrate Synthase Activity—Citrate synthase activity was assayed as described previously with modifications (25). 200 µl of reaction mixture (125 mM Tris-HCl, pH 8, 0.3 mM acetyl-CoA, 0.1% (v/v) Triton X-100, 0.1 mm 5’,5’-dithiobis(2-nitrobenzoic acid)) was incubated with 5 µg of skeletal muscle protein lysate or 10 µg of isolated skeletal muscle mitochondrial protein at 30 °C for 10 min. The reaction was initiated by the addition of freshly prepared 0.5 mM oxaloacetate, and the reduction of 5’,5’-dithiobis(2-nitrobenzoic acid) was continuously monitored by absorbance at 412 nm for 3 min.

NAD Metabolite Extraction from Muscle and Mitochondria—NAD, NAM, and NMN were extracted from ~50 mg of muscle tissue in 0.6 M perchloric acid at 4 °C using a Tissue-Lyzer (Qiagen) set to 20 Hz for 6 min. Lysates were pelleted at 20,000 × g for 10 min at 4 °C and diluted 1:200 in ice-cold 100 mM phosphate buffer, pH 8. For determination of NMN, the supernatant was further neutralized with 1 M potassium carbonate and centrifuged to remove insoluble material. NADH was similarly extracted from ~50 mg of muscle tissue in alkaline extraction buffer (25 mM NH₄Ac, 25 mM NaOH, 50% (v/v) acetonitrile) flushed with nitrogen gas. For measurement by cycling assay, alkaline lysates were mixed 1:1 (v/v) with ethanol extraction buffer (250 mM KOH, 50% (v/v) EtOH) and heated at 55 °C for 10 min. Lysate supernatants were diluted 1:50 in ice-cold 100 mM phosphate buffer, pH 8. NADH was extracted from 100 µg of isolated mitochondria by vortexing vigorously in 100 µl of 0.6 M perchloric.

NAD and NADH Measurement—NAD and NADH were immediately measured after extraction by an enzymatic cycling assay in a 96-well format as described previously with modifications (26). Briefly, 5 µl of NAD standards or diluted tissue extracts was combined with 95 µl of cycling mixture (2% ethanol, 100 µg/ml alcohol dehydrogenase, 10 µg/ml diaphorase, 20 µM resazurin, 10 µM flavin mononucleotide, 10 mM nicotinamide, 0.1% BSA in 100 mM phosphate buffer, pH 8.0). The cycling reaction proceeded for 30 min at room temperature, and resorufin accumulation was measured by fluorescence excitation at 544 nm and emission at 590 nm. NAD and NADH values were confirmed by HPLC analysis of the same extracts.

NAD Metabolite Measurement by HPLC—Separation of NAD, NAM, and NMH was performed on non-diluted extracts as described (27) with modifications using an Adsorbosphere XL ODS column (5 µm, 4.6 × 250 mm) preceded by a guard column at 50 °C. Flow rate was set at 0.4 ml/min. The mobile phase was initially 100% mobile phase A (0.1 M potassium phosphate buffer, pH 6.0 containing 3.75% methanol). Methanol was linearly increased with mobile phase B (0.1 M potassium phosphate buffer, pH 6.0 containing 30% methanol) to 50% over 15 min. The column was washed after each separation by increasing mobile phase B to 100% for 5 min. UV absorbance was monitored at 260 and 340 nm with a Shimadzu SPD-M20A. Separation of NMN was carried out on an YMC-Pack ODS-A column (5 µm, 4.6 × 250 mm) at 30 °C. Flow rate was set at 0.4 ml/min. The mobile phase was initially 100% mobile phase A (0.1 M potassium phosphate buffer, pH 6.0) for the first 8 min. Methanol was linearly increased with mobile phase B (0.1 M potassium phosphate buffer, pH 6.0 containing 30% methanol) to 50% over 7 min. The column was washed after each separation by increasing mobile phase B to 100% for 3 min, and UV absorbance was similarly monitored. Pertinent peak areas were integrated by the LabSolution software from Shimadzu, quantified using standard curves, and normalized to weights of frozen tissues.

Mitochondrial Isolation from Skeletal Muscle—Mice were euthanized by cervical dislocation, and their hind limb muscles dissected and placed immediately in ice-cold muscle homogenization buffer (50 mM Tris-HCl, pH 7.4, 100 mM KCl, 5 mM MgCl₂, 1 mM EDTA, pH 8.0, 18 mM ATP) at pH 7.2. The entire procedure was performed at 4 °C. The fat and connective tissues were removed, and minced muscles were incubated for 2 min in 2.5 ml of homogenization buffer supplemented with 60 units/ml protease from Bacillus licheniformis (Sigma). Muscle pieces were washed twice with 5 ml of unsupplemented homogenization buffer and subjected to an ice-cold motorized homogenizer for 10 min at 150 rpm. A small aliquot of the homogenate was removed and stored at −80 °C for further analysis. The remaining homogenate was centrifuged at 720 × g for 5 min at 4 °C. The pellet was resuspended in homogenization buffer and centrifuged for an additional 5 min at 720 × g. The supernatants were combined and centrifuged at 10,000 × g for 20 min at 4 °C. The supernatant was discarded, and the pellet was resuspended in homogenization buffer and further centrifuged for 10 min at 10,000 × g. The final mitochondrial pellet was resuspended in resuspension buffer (225 mM sucrose, 44 mM KH₂PO₄, 12.5 mM MgOAc, 6 mM EDTA, pH 7.4) at 4 µl of buffer/mg of muscle tissue.

Mitochondrial Oxygen Consumption—Oxygen consumption was measured using the Oroboros High-Resolution Respirometry System (Innsbruck, Austria). Two 2-ml chambers were loaded with 150 µg of isolated skeletal muscle mitochondrial protein in 37 °C MiRO5 respiration medium (110 mM sucrose, 20 mM HEPES, 10 mM KH₂PO₄, 20 mM taurine, 60 mM potassium lactobionate, 3 mM MgCl₂, 6H₂O, 0.5 mM EGTA, 1 g/liter defatted BSA, pH 7.1). 10 mM pyruvate and 5 mM malate or 20 µM palmitoyl carnitine and 5 mM malate were added as oxidative substrates followed by 4 µmol of ADP to stimulate maximal respiration. Glutamate and cytochrome c were added to replenish TCA cycle intermediates and confirm the integrity of mitochondrial membranes, respectively, following the manufacturer’s instructions. To interrogate complex II (succinate dehydrogenase), 1 µM rotenone and 10 mM succinate were added. The assay was ended by addition of 5 µM antimycin A for calculation of residual oxygen consumption.
Histology and Immunohistochemistry—Histological analysis was performed on 4-μm-thick paraffin sections and deparaffinized in Histoclear (National Diagnostics) followed by rehydration through a graded ethanol series. Selected sections were stained with Gill 3 hematoxylin (Thermo Scientific) and eosin (Sigma). For immunohistochemistry, antigens were retrieved following rehydration by microwave heating in EDTA buffer, pH 7.5 for 10 min followed by cooling for 20 min and blocking of endogenous peroxidase with hydrogen peroxide. Slides were incubated overnight at 4 °C with the primary antibody against slow skeletal myosin heavy chain (Abcam, ab11083) diluted 1:3000. Following washing with PBS, slides were incubated with biotinylated secondary antibody (Vector Laboratories, BA-2001) at 37 °C for 1 h, washed three times with PBS, and incubated with streptavidin-conjugated HRP (Vector Laboratories, PK6101) for 1 h at 37 °C. Staining was detected with diaminobenzidine (Dako) chromogen, and slides were counterstained with hematoxylin.

Treadmill, Voluntary Wheel Running, and Grip Strength—Mice were habituated to the Exer3-6 treadmill system with shock detector (Columbus Instruments, Columbus, OH) by running at 10 m/min for 15 min at 0° incline 1 day before the test. To test exercise tolerance, mice were fasted with access to water for 3 h midday before simultaneously running a protocol described in Fig. 3D of 0–25 m/min at 0° incline. Exhaustion was defined as 50 cumulative electrical stimuli (73 V, 0.97 mA, 1 Hz). Exhausted mice were removed from the treadmill and allowed access to food and water. To assess voluntary running, mice were individually housed and allowed access to a computer-monitored running wheel (Columbus Instruments) continuously for 3 weeks. Data were compiled from the 2nd and 3rd weeks after habituation. Grip strength was measured with a rodent grip meter (TSE, Hamburg, Germany) as described previously (28).

Indirect Calorimetry—VO₂, VCO₂, and activity were measured in individually housed mice using an open circuit calorimeter (Columbus Instruments Comprehensive Lab Animal Monitoring System). Room air was supplied at 0.6 liter/min, and exhaust air was sampled at 1-h intervals for 96 h. The respiratory exchange ratio was calculated as the ratio of VCO₂/VO₂ with lower values indicating oxidation of fatty acids. Activity was monitored by infrared beam interruption. Data were compiled from hours 48 to 96 after habituation.

Serum Non-esterified Fatty Acids and Cholesterol—Plasma samples were collected from the tail veins of awake mice after a 5-h fast. Plasma lipids were analyzed individually by analytical chemistry with an automated Cobas Mira Autoanalyzer (Roche Diagnostics).

Lactate, Pyruvate, and Ketone Measurements—The concentrations of lactate, pyruvate, β-hydroxybutyric acid, and acetoacetate were assayed by GC/MS as their acetylated pentafluorobenzyl derivatives as described (29) with modifications. The derivatized analytes were quantified against an internal standard of β-hydroxypentanoate-d₅. Frozen tissues (~40 mg) were spiked with 20 nmol of β-hydroxypentanoate-d₅ and 20 μmol of NaB₂H₄ and homogenized on ice for 2 min in 0.5 ml of a solvent mixture containing equal parts acetonitrile, methanol, and water as described (30). The cold homogenate was then centrifuged at 3000 × g for 10 min at 4 °C. The supernatants were incubated with 20 μmol of pentafluorobenzyl bromide in acetone solution at 60 °C for 1 h. The pentafluorobenzyl bromide-derivatized analytes were extracted with 1 ml of chloroform, dried with nitrogen gas, and further derivatized by acetylation using 30 μl of acetic anhydride and 30 μl of pyridine for 1 h at 65 °C. GC/MS analysis was conducted on an Agilent 5973 mass spectrometer linked to a Model 6890 gas chromatograph equipped with an autosampler and an Agilent OV-225 capillary column (30 m, 0.32-mm inner diameter). The injection volume was 1 μl with a 30:1 split ratio, and the carrier gas was helium at 2 ml/min. The injector temperature was set at 200 °C, and the transfer line was set at 250 °C. The GC temperature program was as follows: 100 °C for 1 min, increase by 10 °C/min to 230 °C followed by 50 °C/min to 300 °C, and hold for 5 min. Chemical ionization GC/MS was monitored for lactate, pyruvate, β-hydroxybutyric acid, and acetoacetate at m/z ratios of 131, 132, 145, and 146, respectively.

Estimation of Compartmental Free NAD/NADH Ratios—Estimates of NAD/NADH ratios in cytosol and mitochondria were calculated from metabolites presented in Table 1 (extracted from whole tissue) according to the equilibrium constants for NAD-linked lactate dehydrogenase and β-hydroxybutyrate dehydrogenase, respectively, as determined previously (31). Calculations according to the following equations assumed enzymatic equilibrium at 38 °C and pH 7.0 in each compartment.

\[
\frac{[\text{NAD}^+]_{\text{cyto}}}{[\text{NADH}]_{\text{cyto}}} = \frac{[\text{pyruvate}][H^+]}{[L-\text{lactate}][1.11 \times 10^{-11}]} \quad (\text{Eq. 1})
\]

\[
\frac{[\text{NAD}^+]_{\text{mito}}}{[\text{NADH}]_{\text{mito}}} = \frac{[\text{acetoacetate}][H^+]}{[\text{β- hydroxybutyrate}][4.93 \times 10^{-7}]} \quad (\text{Eq. 2})
\]

Statistics—Results were analyzed by two-tailed unpaired Student’s t test or one-way analysis of variance with Bonferroni’s multiple comparisons post hoc test as indicated. p < 0.05 was taken to be significant. Analysis was conducted with GraphPad Prism 5 software.

RESULTS

Generation of mNAMPT Mice—To investigate the role of NAD in muscle, we generated mice carrying an exogenous copy of the Nampt cDNA separated from the CAGGS promoter by a floxed stop cassette. Expression of Cre recombinase under control of the muscle creatine kinase promoter (MCK-Cre) results in excision of the stop cassette and overexpression of Nampt in skeletal and cardiac muscle (mNAMPT mice). An important advantage of targeting muscle is that it avoids complications associated with the reported production of extracellular NAMPT (eNampt/PBEF/Visfatin) by adipose tissue (32), immune cells (33), and hepatocytes (34). Deletion of the stop cassette was detected in ~40% of muscle nuclei using a probe specific to the intact allele (Fig. 1A), consistent with efficient recombination of the genomes contained within myofibers (35). Expression of Nampt mRNA increased ~10- to 20-fold in the quadriceps of animals carrying one (NC) or two (NNC)
copies of the transgene, respectively, whereas expression of the 3’-UTR of endogenous Nampt was unaffected (Fig. 1B). Similar increases in Nampt expression were confirmed in gastrocnemius and the more oxidative soleus muscles (data not shown). Transcripts of enzymes downstream in the NAD salvage pathway were unchanged, although expression of both NMNAT1
and NMNAT3 trended higher with increasing Nampt gene dosage (Fig. 1B). Expression of the transcript for nicotinamide N-methyltransferase, which competes with NAMPT to metabolize NAM, was low in skeletal muscle and unaffected by genotype. In accordance with Nampt transcript levels, we observed a robust increase in NAMPT protein in both skeletal (10-fold) and cardiac (4-fold) muscles of NC mice (n = 11 and 3, respectively) and an ~50% increase in intramuscular NAD content (Fig. 1, C–E). Because only a small number of NNC mice were available, we focused primarily on the NC mice for subsequent experiments.

Histological analysis of hind limb muscles containing primarily glycolytic type IIB (extensor digitorum longus) and oxidative type I and IIA fibers (soleus) showed normal polygonal cross-sectional morphology and diameter as well as equivalent staining for type I myosin heavy chain in NC mice as compared with littermate controls (Fig. 1F). NAD levels in heart were also significantly elevated albeit to a lesser degree than in skeletal muscle, consistent with the activity of MCK-Cre in these tissues (36) and the relative expression of NAMPT (Fig. 1G). Gross examination of myocardium did not reveal any major changes in heart tissue, nor was there a change in organ weight (Fig. 1H).

To test whether NAD generation in mNAMPT mice is limited by NAM availability, we administered 400 mg/kg NAM in the drinking water for 1 week and examined skeletal muscle NAD levels. No further elevation of NAD was observed in mNAMPT mice under these conditions, suggesting that NAM availability does not limit NAMPT activity in muscle under standard dietary conditions (Fig. 1I).

Characterization of Muscle Oxidative Function—Based on earlier work using NAD-boosting strategies in mice, we hypothesized that skeletal muscle of mNAMPT mice would exhibit enhanced oxidative function. To test this, we examined several markers of mitochondrial content in gastrocnemius and quadriceps muscles. We found protein expression of subunits from four different complexes of the electron transport chain to be similar in control and mNAMPT mice (Fig. 2A). We further observed similar levels of citrate synthase activity (Fig. 2B) and mtDNA content (Fig. 2C) between the two groups. Consistent with these findings, the Nampt transgene had no significant effect on the expression of genes associated with mitochondrial biogenesis (TFAM, PGC1α, mfsn2, and Sirt1), oxidative phosphorylation (CytC, Cox5b, and Ndufs8), coupling (UCP2), or substrate selection (PDK4, MCAD, and Sirt3) in quadriceps or soleus muscles (Fig. 2E).

Interventions that alter NAD in cultured cells or the liver have been reported to concomitantly alter the mitochondrial NAD pool (10, 20), which may directly influence oxidative...
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We thus hypothesized that mitochondria isolated from the muscles of mNAMPT mice would exhibit an increase in NAD content proportional to the effect in whole muscle tissue. Surprisingly, we found that the NAD content of muscle mitochondria was similar between groups (Fig. 2D). Accordingly, mitochondria did not exhibit any alterations in oxidative capacity that persisted through isolation as detected using high resolution respirometry (Fig. 2, F and G). Mitochondria isolated from control and mNAMPT muscle showed similar patterns of oxygen consumption across all complexes on both pyruvate and fatty acid substrates. These results suggest that NAMPT overexpression is not sufficient to promote the biogenesis or respiratory capacity of muscle mitochondria.

We next sought to test the influence of NAMPT on muscle performance in vivo. We found voluntary ambulatory and wheel running activity across the light/dark cycle to be indistinguishable between control and NC mice (Fig. 3, A and B). Forelimb strength and treadmill running capacity were also comparable (Fig. 3, C–E). Together, these data indicate that NAD is unlikely to limit muscle metabolic function in healthy wild type mice.

Muscle NAMPT Does Not Protect against Metabolic Consequences of High Fat Diet—Genetic and pharmacological interventions that elevate NAD levels have been reported to ameliorate many of the negative consequences of high fat diet feeding, including insulin resistance and weight gain. To investigate whether NAMPT-mediated NAD synthesis in muscle conferred similar protective effects, we placed mNAMPT mice on an HFD for 24 weeks. No significant differences in weight gain, plasma free fatty acids, or total cholesterol were observed between control and NC mice (Fig. 4, A–C). After 10 weeks on respective diets, HFD-fed NC and control mice exhibited similar declines in glucose tolerance, VO₂, and respiratory exchange ratio as compared with chow-fed littermates (Fig. 4, D, E, and G–J). Interestingly, HFD caused a slight but significant decline in intramuscular NAD in control mice that was completely rescued by the transgene (Fig. 4F). Thus, intramuscular NAD levels do not appear to have a major influence on the general metabolic deterioration caused by HFD.

NAMPT Overexpression in Muscle Does Not Significantly Affect Steady-state Levels of NAM or NMN—NAM, a substrate of NAMPT, is also a non-competitive inhibitor of sirtuins and a competitive inhibitor of PARPs. Thus, one mechanism by which NAMPT overexpression might influence such enzymes is through the depletion of NAM rather than through the generation of NAD per se. Contrary to our initial expectation, we did not observe a significant depletion of NAM in quadriceps muscles of mNAMPT mice (Fig. 5A), potentially indicating that this metabolite is rapidly replenished. We also found no significant accumulation of NMN, the direct product of NAMPT, in mNAMPT muscle (Fig. 5B), supporting the view that NAMPT catalyzes the rate-limiting step in the NAD salvage pathway.

Muscle Redox State Is Independent of NAMPT-mediated NAD Synthesis—The reduced form of NAD, NADH, acts as a key electron donor and allosteric regulator of mitochondrial dehydrogenases, and the ratio of oxidized to reduced nucleotides can influence the rates of many reactions within the cell. Although overexpression of NAMPT in cells and treatment of mice with NR have both been reported to increase the NAD/
NADH redox ratio (20, 23, 38), it remains unresolved whether changes in absolute NAD concentration or the redox ratio play a greater role in mediating the effects of NAD-boosting interventions. Interestingly, total concentrations of NADH were significantly elevated in the muscles of mNAMPT mice, reflecting an apparent equilibration between the oxidized and reduced forms of NAD to maintain the NAD/NADH ratio across a range of absolute concentrations (Fig. 5, C and D).

FIGURE 4. Overexpression of NAMPT in muscle does not protect against metabolic consequences of high fat diet. A, body weights of mice provided a standard chow diet (CD) or HFD for 24 weeks starting at 3 months of age (n = 6 mice per group). B and C, plasma non-esterified fatty acids (NEFA) and total cholesterol after 24 weeks on the indicated diets (n = 7–10 per group). D and E, glucose tolerance after 20 weeks on the indicated diets (n = 6 per group). AUC, area under the curve. F, NAD content of gastrocnemius muscles after 24 weeks on the indicated diets (n = 5–6 per group). G, oxygen consumption (VO₂) normalized to body weight after 24 weeks on the indicated diets. The time course represents the 48-h period following habituation. Shaded panels indicate periods of darkness (n = 5 per group). H, average VO₂ during light and dark periods shown in G. I, respiratory exchange ratios (RER) over the same time course shown in G. J, average respiratory exchange ratio during light and dark periods shown in I. Error bars represent S.E. *, p < 0.05; **, p < 0.01; ns, not significant by Student’s t test.
A large fraction of the intracellular NADH pool is protein-bound and therefore inaccessible to enzymatic reactions that would lead to its oxidation. NAD(H) is further compartmentalized into mitochondrial and nucleocytosolic pools, suggesting that significant changes in the abundance of free nucleotides within a given compartment could be missed by measuring only total tissue levels. To confirm the maintenance of redox state in the nucleocytosolic compartment of mNAMPT muscle, we...
TABLE 1
Concentrations of redox-sensitive metabolites in quadriceps muscle of 5-month-old mice

| Group     | Observations | Lactate | Pyruvate | \( \beta \)-HB | Acetoacetate |
|-----------|--------------|---------|----------|----------------|--------------|
| Control   | 7            | 14.3 (1.1) | 0.289 (0.029) | 0.086 (0.007) | 0.131 (0.015) |
| NC        | 6            | 12.7 (0.9)  | 0.313 (0.042) | 0.058 (0.005) | 0.133 (0.020) |
| NNC       | 3            | 12.2 (1.7)  | 0.285 (0.042) | 0.058 (0.006) | 0.118 (0.015) |

Although obese control animals exhibited diminished metabolic health coincident with a small but significant decline in skeletal muscle NAD, this decline was completely rescued by the Nampt transgene to no effect (Fig. 4F). Thus, the slight decrease in muscle NAD does not contribute meaningfully to the detrimental consequences of obesity, nor does elevating NAD have a protective effect.

Although little is known about the dynamics of pyridine nucleotide transport across the mitochondrial membranes, it is appreciated that mitochondria maintain bioenergetics via a distinct NAD pool that is reduced to NADH in a much higher proportion than the nucleocytosolic pool (31) and is shielded from the activity of nuclear and cytosolic enzymes (10, 43, 44). Although distinct, the mitochondrial NAD pool tends to mirror oxidative metabolism in the liver where NAD deficiency can limit mitochondrial oxidative metabolism (20, 37). Although we observed a slight trend toward an increase in mitochondrial NAD content in muscle (Fig. 2D), the effect was clearly dampened compared with the increase in whole tissue. Moreover, mitochondria isolated from mNAMPT mouse consumed pyruvate and palmitoyl carnitine at rates equivalent to those of control littermates (Fig. 2G), consistent with the VO2 and respiratory exchange ratio measurements from whole animals (Fig. 4, H and J).

DISCUSSION

Chronic deficiency of dietary NAD precursors is known to cause the progressive and lethal disease pellagra (39), but the more recent notion that increasing NAD availability might suffice to enhance the metabolic fitness of an otherwise healthy tissue is just beginning to be explored. We report that targeting the NAD salvage pathway in muscle via overexpression of NAMPT increases the NAD content of skeletal muscles by ~50% but does not overtly influence muscle physiology or whole body energy balance in the context of standard or high fat diets. Our findings represent the first investigation of these phenotypes using a tissue-specific approach to enhance NAD synthesis and in stark contrast to the beneficial effects that have been observed using a variety of genetic and pharmacological approaches to elevate NAD throughout the whole body.

It is noteworthy that our model achieved a relative elevation of skeletal muscle NAD greater than or equal to those reported for small molecule precursor supplementation (20, 40, exercise (3, 4), or deletion of either PARP-1 (18) or PARP-2 (19). Because increased NAD was achieved via supraphysiological levels of NAMPT protein expression and was not further enhanced by supplementing NAM, it is possible that negative feedback maintains the steady-state NAD concentration in our model at or near a theoretical limit for salvage-mediated synthesis. These observations are consistent with reports that the \( K_m \) of NAMPT for NAM in the presence of ATP is well below our observed tissue concentrations of NAM and that physiological levels of pyridine nucleotides can inhibit NAMPT activity in rat liver extracts (41, 42). Despite the elevated steady-state concentration of muscle NAD that was achieved, mNAMPT mice failed to display phenotypes suggesting any alteration of mitochondrial content (Fig. 2, A–C) or transcriptional adaptations (Fig. 2E), consistent with their unchanged strength and exercise capacity (Fig. 3, B–E). In addition, mNAMPT mice gained weight normally on a high fat diet and did not display changes in glucose tolerance or oxygen consumption (Fig. 4, E and H).

3 D. W. Frederick, A. Dávila, Jr., and J. A. Baur, unpublished results.
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els as was observed in the context of PARP-1 deletion (18). Our results may indicate that (a) flux through the salvage pathway is unchanged, (b) increased NAM consumption is counterbalanced by a corresponding increase in NAM generation by NAD-dependent enzymes, or (c) NAM exchanges continuously with the plasma to maintain a stable tissue concentration irrespective of NAMPT activity. The cellular NAD/NADH ratio is sensitive to metabolic flux and extrinsic factors and can modify enzymatic activity independently from changes in NAD concentration per se (38, 47). For example, the transcriptional co-repressor C-terminal binding protein is reciprocally regulated by the oxidized and reduced nucleotides (48), and it is debated whether the sirtuins are competitively inhibited by NADH in vivo (14, 46, 47). Thus, any changes in the NAD/NADH ratio could mediate effects of NAD-boosting interventions. Contrary to our expectation that NAMPT overexpression would increase the skeletal muscle NAD/NADH ratio as reported in isolated myoblasts (38) and heart (11) as well as muscles of NR-fed mice (20), we observed equilibration between the oxidized and reduced nucleotides to maintain the redox state in mNAMPT muscle (Fig. 5D). Several key differences distinguish our experiment from previous studies. First, experiments in cultured myoblasts may not accurately model the changes in nutrient availability, glycolytic flux, or ATP demand experienced by muscle fibers in vivo; all of these affect the redox state of NAD. Moreover, the redox states of liver and other tissues may influence muscle in vivo via circulating redox-active metabolites, including lactate, pyruvate, and ketone bodies. Second, in the cardiac model of Hsu et al. (11), NAMPT overexpression shifted the redox ratio in concert with a larger increase in NAD than has been reported for any other intervention (~5-fold), yet a partial equilibration was still apparent. Lastly, the dietary delivery of NR is distinct from our genetic model in that it amounts to a series of discrete systemic administrations, which may be responsible for its effect on muscle redox state. Our finding that redox state was not chronically affected in mNAMPT mice indicates that the metabolic processes responsible for establishing the NAD/NADH ratio are rapid relative to the basal rate of NAD synthesis. Analogous observations were made following deletion of the circadian regulator Rev-erbα in which the muscle NAD/NADH ratio was maintained in the context of a diminished NAD pool (50). Our data indicating that lactate/pyruvate and β-hydroxybutyrate/acetocacetate ratios were unchanged in the muscles of transgenic mice (Fig. 5, G and H) provide additional evidence that NAD equilibrates with NADH and, insofar as the associated enzymes can be presumed to be in equilibrium, adds compartment-specific resolution to both nucleocytosolic and mitochondrial pools of the free nucleotides. Thus, homeostatic mechanisms appear to maintain the NAD/NADH ratio in muscle despite a chronic 50% increase in the size of the NAD pool.

It is interesting to consider why increased NAD availability did not have clear effects on skeletal muscle physiology. Unlike yeast where sirtuins are the most prominent consumers of NAD (51), mammalian genomes encode several additional classes of NAD-consuming enzymes that play important roles in signaling and metabolism and depend on the same NAD pool (52). The evolution of different classes of NAD-dependent enzymes was likely accompanied by additional layers of regulation responsible for modifying the enzyme activities in different contexts. In support of this view, the best studied mammalian sirtuin, SIRT1, is now appreciated to have numerous binding partners and post-translational modifications (53) at least some of which can regulate deacetylase activity independently from changes in NAD (54). Therefore, it will be important to identify the specific tissues or cell types that are capable of responding directly to changes in NAD availability.

Overall, our findings using a conditional Nampt allele support a model in which the effects of globally increasing NAD availability on muscle physiology are not muscle-autonomous. It remains possible that more profound muscle-autonomous effects would emerge following larger increases in NAD, concurrent with energetic stress, or in a diseased state. However, we note that none of these conditions were required for NR supplementation or PARP-1 deletion to benefit muscle (18, 20). Interestingly, overexpression of the NAD-dependent deacetylase SIRT1 in the brain has recently been shown to improve muscle physiology (55, 56), whereas overexpression of the enzyme in the muscle itself was ineffective (57). Moreover, NMN can promote insulin secretion in cultured islets and in vivo, consistent with the stimulation of SIRT1 in pancreatic β cells (7, 21), and knockdown of nicotinamide–N-methyltransferase was recently shown to elevate adipose tissue NAD levels in concert with increased whole body energy expenditure (49). Thus, it is possible that improvements in muscle physiology during NAD precursor supplementation occur via pleiotropic effects of NAD-dependent enzymes in tissues other than skeletal muscle. It will be important to determine in future studies whether the central nervous system, hormones, circulating metabolites, or a combination thereof mediates the effects of global NAD availability on muscle physiology. It will also be interesting to determine the tissue-autonomous impact of enhanced NAD salvage capacity under conditions known to deplete the metabolite, including genotoxic stress (10) and the course of natural aging (21, 40).

In summary, our muscle-specific transgenic approach provides an important step toward dissecting the mechanisms by which NAD availability influences mammalian physiology. A deeper understanding of the tissues and signals that mediate these effects will greatly facilitate the development of NAD-boosting therapeutics to combat comorbidities of obesity and aging.

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