NADH inhibition of SIRT1 links energy state to transcription during time-restricted feeding

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In mammals, circadian rhythms are entrained to the light cycle and drive daily oscillations in levels of NAD+, a cosubstrate of the class III histone deacetylase sirtuin 1 (SIRT1) that associates with clock transcription factors. Although NAD+ also participates in redox reactions, the extent to which NAD(H) couples nutrient state with circadian transcriptional cycles remains unknown. Here we show that nocturnal animals subjected to time-restricted feeding of a calorie-restricted diet (TRF-CR) only during night-time display reduced body temperature and elevated hepatic NADH during daytime. Genetic uncoupling of nutrient state from NADH redox state through transduction of the water-forming NADH oxidase from Lactobacillus brevis (LbNOX) increases daytime body temperature and blood and liver acyl-carnitines. LbNOX expression in TRF-CR mice induces oxidative gene networks controlled by brain and muscle Arnt-like protein 1 (BMAL1) and peroxisome proliferator-activated receptor alpha (PPARα) and suppresses amino acid catabolic pathways. Enzymatic analyses reveal that NADH inhibits SIRT1 in vitro, corresponding with reduced deacetylation of SIRT1 substrates during TRF-CR in vivo. Remarkably, Sirt1 liver nullizygous animals subjected to TRF-CR display persistent hypothermia even when NADH is oxidized by LbNOX. Our findings reveal that the hepatic NADH cycle links nutrient state to whole-body energetics through the rhythmic regulation of SIRT1.

Metabolic homeostasis in mammals is mediated by interlocked nutrient-sensing and temporal signals throughout the 24-h light/dark cycle. The molecular clock network drives oscillations of a broad range of transcripts and metabolites that direct anabolic and catabolic metabolism in anticipation of the fasting/feeding–sleep/wake cycle1–3. Interactions between the core molecular clock and nutrient-responsive transcription factors (TFs) contribute to metabolic homeostasis, yet how these pathways cooperate under long-term energy-deficient conditions has remained obscure.

In yeast, calorie restriction (CR) transcriptionally reprogrammes metabolic gene expression to shift oxidative fuel preference and maintain energetic homeostasis of the cell4. In mammals, CR downregulates energy-intensive processes, such as thermogenesis during sleep, and upregulates anabolic processes within the liver that convert metabolite stores into energetic substrates for the brain5–8. NAD+ and the NAD+-dependent sSir2p deacetylase are required for the transcriptional response to CR in yeast, although the role of NAD+ in the response to low-energy conditions in mammals remains less clear.

Transcriptomic studies have revealed robust oscillations in the expression of the rate-limiting enzyme involved in NAD+ biosynthesis across peripheral tissues, which in turn feedback to regulate metabolic transcription cycles through SIRT1-mediated deacetylation and inhibition of the circadian repressor PER2 (refs. 9–11). Supplementation of NAD+ with the soluble precursor nicotinamide riboside in ageing mice reverts senescence of the sleep/wake and mitochondrial oxidative respiration cycles, suggesting that robustness of the NAD+-SIRT1-clock pathway may enhance fitness12. NAD+ also functions as an electron shuttle in oxidoreductase equilibrium reactions that vary according to both nutrient availability and time of day13,14, yet whether metabolic control of rhythmic transcription is modulated by cyclic changes in NAD(H) balance remains unknown.

To investigate whether NAD(H) balance provides a signal to regulate transcriptional and metabolic cycles in vivo, we examined the effect of uncoupling nutrient state from NADH levels by tonically inducing NADH oxidation during CR through hepatic transduction of LbNOX15,16. When NADH levels were dissociated from energy state using LbNOX, we observed transcriptional reprograming of SIRT1- and BMAL1-mediated gene networks that regulate fatty acid and amino acid metabolism and thermogenesis. Our studies reveal that NADH redox state drives energy conservation during sleep through rhythmic inhibition of SIRT1 and downstream circadian processes.

Results
Elevation in hepatic NADH drives morning torpor with TRF-CR. We first sought to examine how changes in energy state during a TRF-CR diet impact NAD+ and NADH across the fasting/feeding cycle compared with time-restricted feeding of a regular chow
Liver is a key metabolic organ that governs whole-body metabolism during fasting and low-nutrient conditions by releasing metabolites into the circulation that can be oxidized by other tissues, including brain and brown adipose tissue (BAT)\(^21\). For example, the partial oxidation of fatty acids (derived from white adipose tissue during fasting/sleep) in the liver produces acyl-carnitines that are secreted into serum and serve as a thermogenic fuel source for BAT\(^24-26\). To determine whether TRF-CR alters hepatic mobilization of metabolite stores during the daytime, we first profiled acyl-carnitines in both liver and serum from TRF-CR mice using tandem mass spectrometry (MS/MS), as described previously\(^27,28\). TRF-CR decreased acyl-carnitine species of all lengths in serum (Fig. 1c), and medium- and long-chain acyl-carnitine species in liver (Extended Data Fig. 1g), suggesting that hepatic production of acyl-carnitines was reduced with TRF-CR. Because hepatic acyl-carnitine production has been linked to regulation of body temperature at the level of BAT\(^14\), we next examined body temperature rhythms by implanting temperature transponders into freely moving mice during TRF-CR. As has been reported, mice on regular chow diet demonstrate a circadian variation in body temperature (Fig. 1d)\(^29,30\). TRF-CR led to a daily rhythmic reduction in core body temperature relative to TRF-CR mice, predominantly during the early daytime hours (Fig. 1d), consistent with previous reports in one-meal CR paradigms\(^31\). We note that the body temperature of TRF-CR mice is not directly dependent upon the availability of food pellets, because body temperature begins to: (1) decrease in the late night-time before access to the last pellet of food (at ZT22); and (2) normalize at midday (at ZT8), 4 h before refeeding, when animals are still fasting. Together, these results reveal that TRF-CR leads to elevation of hepatic NADH content concurrent with reduced acyl-carnitine production and the nadir in daily core body temperature rhythms during CR.

To determine whether the increase in NADH per se inhibits hepatic lipid oxidation during TRF-CR, we took advantage of a novel genetic tool recently developed to manipulate NADH levels without changing the nutrient state of the animal using flavin adenine dinucleotide-dependent \(Lb\)NOX, which reduces oxygen to water and has specificity for oxidizing NADH over NADPH (Fig. 1e)\(^16,17\). Although NADH exists in distinct mitochondrial and cytonuclear pools, we chose to utilize the cytonuclear form of \(Lb\)NOX to manipulate NADH levels because the increased lactate flux into gluconeogenesis during TRF-CR (Extended Data Fig. 1d) suggests a net increase in cytosolic NADH due to the export of NADH equivalents from the mitochondria into the cytonuclear compartment via the malate–aspartate shuttle\(^10-12\), although accurate subcellular quantification is not possible in liver. Before subjecting mice to TRF-CR, we transduced cytonuclear \(Lb\)NOX under a liver-specific promoter to wild-type mice by retro-orbital injection of adeno-associated viruses (AAVs) that expressed either \(Lb\)NOX or an empty vector (‘null’). We observed liver-specific \(Lb\)NOX expression (Fig. 1e) and reduced levels of daytime NADH during TRF-CR without changes in the levels of oxidized NAD\(^+\) (Fig. 1f). To determine whether the altered acyl-carnitine profiles and whole-body temperature rhythms during TRF-CR depend on elevated hepatic NADH during TRF-CR, we assessed acyl-carnitine levels in liver and serum.

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**Fig. 1 | Elevated NADH drives the daytime dip in body temperature during time-restricted calorie restriction.** a, Model for the TRF-CR diet using an automated feeder system in 4-6-month-old male C57BL/6J mice. Control mice received a 300 mg pellet of regular chow every 1.2 h throughout the dark period (TRF-Reg), whereas TRF-CR mice received a 300 mg pellet of a carbohydrate-depleted, nutrient-controlled chow every 2 h throughout the dark period, resulting in a 40% reduction in calories. b, Relative concentration of NADH and NAD\(^+\) by HPLC in liver of TRF-Reg \((n=12)\) and TRF-CR \((n=11)\) mice during the daytime (ZT4) (Fig. 1b), whereas TRF-CR mice received a 300 mg pellet of a carbohydrate-depleted, nutrient-controlled chow every 2 h throughout the dark period, resulting in a 40% reduction in calories. Relative concentration of NADH and NAD\(^+\) by HPLC in liver of TRF-Reg \((n=12)\) and TRF-CR \((n=11)\) mice during the daytime (ZT4). c, \(\log_2\) (fold change) in daytime serum acyl-carnitine levels during TRF-CR compared with TRF-Reg mice \((n=3)\). d, Body temperature rhythms monitored non-invasively using subcutaneous probes in TRF-Reg \((n=12)\) and TRF-CR \((n=9)\) mice over 24 h (double plotted for clarity). e, Model depicting NADH-consuming reaction of \(Lb\)NOX. Representative tissue-specific expression profile of cytonuclear \(Lb\)NOX in null- and \(Lb\)NOX-transduced mice relative to \(Lb\)NOX-transduced liver. f, Relative concentration of NADH and NAD\(^+\) in liver of null- and \(Lb\)NOX-transduced TRF-Reg \((n=6)\) or TRF-CR \((n=5)\) for NADH, \(n=6\) for NAD\(^+\) mice during the daytime. g, \(\log_2\) (fold change) in daytime serum acyl-carnitine levels in TRF-CR mice transduced with cytonuclear \(Lb\)NOX compared with TRF-CR mice transduced with null virus \((n=3)\). h, Body temperature rhythms in \(Lb\)NOX-expressing TRF-Reg \((n=12)\) or TRF-CR \((n=9)\) mice over 24 h (double plotted for clarity). Data are presented as mean ± s.e.m. Statistics were performed with unpaired, two-tailed Student’s t test unless otherwise noted in the figure. *\(P<0.05\), **\(P<0.001\). ANOVA, analysis of variance; Cereb, cerebellum; DC, dicarboxylate; Gastroc, gastrocnemius muscle; gWAT, gonadal white adipose tissue; Hypoth, hypothalamus; i, isomer; iWAT, inguinal white adipose tissue; OH, hydroxy; Quad, quadriceps muscle.
from \textit{LbNOX}-expressing TRF-CR-fed mice following monitoring of body temperature rhythms from implanted temperature transponders. Remarkably, we observed that hepatic \textit{LbNOX} overexpression resulted in increased acyl-carnitine levels in both liver (Extended Data Fig. 1h) and serum (Fig. 1g) during TRF-CR and abrogated the daily reduction in core body temperature characteristic of the

\begin{itemize}
  \item \textbf{a} Experimental conditions for time-restricted calorie restriction
  \begin{tabular}{|c|c|}
    \hline
    Light (day) & Dark (night) \\
    \hline
    No food & Food at even intervals \\
    \hline
  \end{tabular}
  \begin{itemize}
    \item Regular chow (TRF-Reg)
    \item Calorie restriction (TRF-CR)
  \end{itemize}

  Outputs:
  \begin{itemize}
    \item Hepatic metabolism
    \item Body temperature
    \item Transcription
  \end{itemize}

  \textbf{b} TRF-CR increases hepatic NADH in the daytime

  \textbf{c} TRF-CR decreases serum acyl-carnitines in the daytime

  \textbf{d} TRF-CR decreases body temperature in the daytime

  \textbf{e} Liver-specific expression of cytonuclear \textit{LbNOX}

  \textbf{f} \textit{LbNOX} reduces daytime NADH during TRF-CR

  \textbf{g} \textit{LbNOX} increases daytime serum acyl-carnitines during TRF-CR

  \textbf{h} TRF-CR does not decrease daytime body temperature in \textit{LbNOX} mice

\end{itemize}
TRF-CR mice (Fig. 1b). These results indicate that the elevated levels of cytonuclear NADH in the liver during daytime both inhibits acyl-carnitine production and decreases core body temperature during TRF-CR.

NADH regulates metabolic transcription factors during TRF-CR. NAD⁺ supplementation increases core clock transcriptional activity and reprogrammes genome-wide transcription of oxidative genes collaboratively controlled through BMAL1–PPARα/β. Further, in yeast, the transcriptional response to CR requires rate-limiting enzymes in NAD⁺ biosynthesis. Thus, we next sought to understand whether the changes in NADH that occur in the cytonuclear compartment of mouse liver during TRF-CR affect genome-wide transcription of metabolic and oxidative gene networks. We performed RNA sequencing (RNA-Seq) in liver of null- and LbNOX-transduced TRF-CR and TRF-Reg mice during the daytime (Fig. 2a and Extended Data Fig. 2a). We identified 930 genes differentially expressed by TRF-CR versus TRF-Reg (DESeq2, false discovery rate (FDR)-adjusted P < 0.05) (Fig. 2a, left). Of those 930 differentially expressed genes, only 57 were still differentially expressed by TRF-CR in the presence of LbNOX (Fig. 2a, middle), whereas 19 genes were uniquely differentially expressed by TRF-CR in the presence of LbNOX (Fig. 2a, right). We next examined the effect of LbNOX on the expression of the 930 genes regulated by TRF-CR. We observed that LbNOX expression in TRF-CR mice caused opposite changes in expression relative to TRF-CR for 95% of genes (Fig. 2b) (binomial test P < 2.2 × 10⁻16). These observations indicate that reduction of NADH levels by LbNOX counteracts the transcriptional response to TRF-CR for a majority of genes that are differentially expressed by TRF-CR during the daytime. Of note, principal component analysis revealed that LbNOX did not affect gene expression in the night (ZT16) (Extended Data Fig. 2c), suggesting that LbNOX regulates the transcriptomic response to TRF-CR specifically in the daytime (ZT4) when NADH is elevated.

We next applied unbiased in silico transcriptomic and motif analyses to identify the gene networks and TFs that were altered by LbNOX during TRF-CR. Using gene ontology analysis, we found that genes that were decreased by TRF-CR and increased by LbNOX (green—group 1) were enriched in gene ontologies associated with fatty acid oxidation and PPARα signalling, whereas genes that were increased during TRF-CR and decreased by LbNOX (red—group 2) were enriched in gene ontologies associated with amino acid metabolism (Fig. 2c). We note that although we found gene ontologies associated with fatty acid and amino acid metabolism, we did not observe enrichment of gene ontologies associated with either glutoneogenesis or glycolysis (Extended Data Fig. 2b). The discovery of gene ontologies associated with fatty acid metabolism is consistent with our finding that TRF-CR decreases acyl-carnitine levels through a mechanism that is reversed by LbNOX (Fig. 1, Extended Data Fig. 1).

Using DNA motif analysis, we next found that canonical PPARα-regulated genes were specifically enriched among those genes that were upregulated in LbNOX-expressing TRF-CR mice (group 1 above), whereas we found enrichment of the bHLH DNA motif that is recognized by CLOCK/BMAL1 among genes that were either up- or downregulated by LbNOX in TRF-CR mice (groups 1 and 2) (Fig. 2d). Further, the hepatic transcriptome pattern of mice subjected to TRF-CR has a strong positive correlation with the hepatic gene expression profile of Ppara nulligous mice (Fig. 2e, top) (log₂(fold change) cut-off ± 0.5; 76% of genes, binomial test P < 7.346 × 10⁻16). The observation that patterns of NADH-sensitive gene expression in liver of TRF-CR mice overlap with that of PPARα-null animals indicates a role for redox state in the transcription control of acyl-carnitine production and thermogenesis.

Interestingly, we observed that Bmal1 ablation is sufficient to decrease expression of Ppara and other fatty acid oxidation genes that are decreased by TRF-CR and increased by LbNOX (Extended Data Fig. 2d), consistent with previous findings that BMAL1 is a positive regulator of fatty acid oxidation and PPARα expression. Comparison of the genome-wide expression profile of TRF-CR with that of Bmal1 ablation (GSE133989) revealed a strong positive correlation in gene expression between these conditions (log₂(fold change) cut-off ±0.5; 70% of genes, binomial test P < 1.17 × 10⁻11) (Fig. 2e, bottom). To test whether hepatic BMAL1 also plays a role in the thermogenic response to lower-nutrient conditions, we measured body temperature in mice harbouring a floxed allele of Bmal1 before and after administration of AAV8-thyroid binding globulin-improved Cre recombinase (AAV8-TBG–iCre) to specifically delete Bmal1 in the liver. When mice were fostered at the start of the light period (ZTO) to induce a low-energy state, we found that liver-specific Bmal1 knockout animals displayed decreased body temperature (Extended Data Fig. 2e).

Hepatic NADH controls amino acid metabolism during TRF-CR. Because gene networks involved in amino acid metabolism were increased in expression by TRF-CR and decreased by LbNOX (Fig. 2c, red), we next measured concentrations of amino acid and hydrophilic metabolites using liquid chromatography–mass spectroscopy (LC–MS) in liver isolated from null- or LbNOX-transduced mice maintained on TRF-Reg and TRF-CR diets. We observed that amino acids involved in nitrogen metabolism (including lysine, asparagine, glutamine and aspartate) were decreased by TRF-CR and increased by LbNOX (Fig. 2f and Supplementary Table 2) (multiple t test, FDR-adjusted P < 0.05). This finding is consistent with enrichment of gene ontology pathways associated with the urea cycle, which sequesters nitrogen (Fig. 2c). Interestingly, we
also observed that oxygen/sulfur-containing amino acids such as serine and methionine were decreased by TRF-CR and increased by LbNOX (Fig. 2f and Supplementary Table 2) (multiple t test, FDR-adjusted P < 0.05), consistent with the enrichment of gene ontology terms associated with cysteine and methionine metabolism (Fig. 2c). Specifically, we observed increased expression of RNAs encoding enzymes within the methionine metabolic pathway, such as Mat1a, Ahcy, Cbs and Cth, in TRF-CR mice, and decreased expression of these genes in LbNOX-expressing mice (Extended Data Fig. 2f). We similarly observed that Bma1 nullizygous animals exhibited increased expression of Mat1a, Cbs and Cth (Extended Data Fig. 2f). To interrogate the extent to which metabolites downstream of methionine and serine are regulated by NADH during TRF-CR, we quantified metabolites by LC–MS in liver and serum from null virus and LbNOX TRF-CR mice. Consistent with the transcriptional analyses, our metabolomic stud-
Fig. 3 | NADH inhibits SIRT1 in the morning during TRF-CR to regulate metabolism and body temperature. a, Model to examine the role of SIRT1 in the NADH-dependent effects on RNAs, metabolites and body temperature during TRF-CR. b, Deacetylation rate for SIRT1 with increasing concentrations of NAD⁺ and NADH assayed by SAMDI-MS (n = 4). c, Model of pyruvate/lactate equilibrium shows that supplementation with pyruvate and lactate reduces and elevates NADH, respectively. Western blotting for SIRT1 targets, Ac-H3K9 and Ac-H4K16, in immortalized mouse embryonic fibroblasts treated with pyruvate or lactate to modulate NADH (n = 3). d, Western blotting for SIRT1 targets, Ac-p53 and Ac-FOXO1, in TRF-Reg or TRF-CR liver of 4–6-month-old male (d) control or liver-specific Sirt1–/– mice or (e) null- or LbNOX-transduced mice. n.s., non-specific. Uncropped western blot scans labelled with LbNOX in TRF-CR, metabolic effects of LbNOX in TRF-CR, SIRT1 required for transcriptional and metabolic effects of LbNOX in TRF-CR, SIRT1 required for body temperature increase from LbNOX during TRF-CR, and for control, n = 5 for L-Sirt1–/– or LbNOX-expressing (n = 5) control and liver-specific Sirt1–/– mice on TRF-CR. Data are presented as mean values ± s.e.m.
ies revealed that catabolites within the methionine/serine pathway, including \(\alpha\)-aminobutyrate and \(\alpha\)-hydroxybutyrate, were increased by TRF-CR and decreased by \(Lb\)NOX in TRF-CR (Extended Data Fig. 2g). We note that within the methionine metabolic pathway, the conversion of \(\alpha\)-ketobutyrate into \(\alpha\)-hydroxybutyrate by LDHA is the only biochemical reaction that utilizes NAD(H) as a cofactor (Extended Data Fig. 2g). These results reveal that NADH redox state during daytime controls both methionine and serine catabolism in response to TRF-CR. Together, these data suggest that elevated daytime NADH during TRF-CR inactivates BMAL1 and PPAR\(\alpha\) in the liver, causing decreased expression of fatty acid oxidation genes, increased expression of genes involved in amino acid metabolism and reduced core body temperature (Fig. 2g).

**NADH inhibits SIRT1 to link energy state to transcription.** Our data suggest that elevated daytime NADH during TRF-CR inactivates BMAL1 and PPAR\(\alpha\) in the liver (Figs. 1 and 2). Because nutrient-dependent deacetylase SIRT1 regulates the activity of both BMAL1 and PPAR\(\alpha\)\(^{31,36,44}\), we hypothesized that the elevated NADH during TRF-CR may inactivate BMAL1 and PPAR\(\alpha\) through SIRT1 (Fig. 3a). The relationship between NADH levels and sirtuin activity in response to time-restricted hypocaloric feeding remains uncertain in mammals, with inconsistent findings from prior biochemical analyses concerning the role of NADH due to differences across isoforms, species and assay conditions\(^{45-47}\). We therefore sought to re-evaluate SIRT1 activity under physiological range concentrations of both NAD\(^+\) and NADH by using a direct MS method, self-assembled monolayers for matrix-assisted laser desorption/ionization (SAMDI-MS)\(^{48}\), which circumvents problems associated with fluorescence-based assays of deacetylase function\(^{49-51}\). Recombinant SIRT1 was incubated with acetyl-peptide substrate (GRK\(^\text{AcRVC}\)) in the presence of varying concentrations of NAD\(^+\) (0–2 mM) and NADH (0–1 mM), followed by quenching and immobilization of the acetylated and deacetylated peptides on a maleimide-functionalized gold surface and quantification of the deacetylation rate by MS. Whereas NAD\(^+\) stimulated SIRT1 deacetylase activity as expected, increasing concentrations of NADH inhibited the activity of SIRT1 (Fig. 3b and Extended Data Fig. 3a). Importantly, the inhibition constant \(K_{\text{NADH}}\) value for SIRT1 that we observed (0.124 mM) falls within previously reported ranges for

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**Fig. 4 | \(Lb\)NOX redox state in liver drives energy conservation during nocturnal CR feeding.** TRF-CR generates rhythmic bouts of daytime torpor through increased levels of NADH in liver, inhibition of SIRT1 and downregulation of oxidative gene networks controlling acyl-carnitines and core body temperature. Reducing levels of NADH in the morning through the transduction of \(Lb\)NOX in TRF-CR mice increases lipid oxidation through the activation of SIRT1, resulting in elevated daytime body temperature rhythms. These findings identify NAD(H) redox state as a link between TRF-CR and whole-body metabolism.
physiological concentrations of total NADH in mouse liver (100–300 pmol per mg liver, or 0.1–0.3 mM)\(^{22,34}\) and is within range of our semiquantitative determination of total NADH levels during TRF-CR (~120 pmol per mg liver, or ~0.12 mM), estimated based on a threefold increase in daytime NADH levels relative to mice fed ad libitum (Extended Data Fig. 1c). Thus, SIRT1 is inhibited by NADH at concentrations that have been reported for total NADH in the whole liver, although accurate determination of NADH in the nucleus of liver is not currently possible.

To test whether NADH inhibits SIRT1 in vivo, we first supplemented mouse embryonic fibroblasts with lactate (to increase NADH) or pyruvate (to decrease NADH) and analysed the acetylation state of the targets of SIRT1 by immunoblotting (Ac-H3K9, Ac-H4K16) (Extended Data Fig. 3b,c)\(^{26,28}\). Acetylated H3K9 and H4K16 increased following lactate supplementation (Extended Data Fig. 3c), suggesting reduced SIRT1 activity. To examine whether elevated daytime NADH levels during TRF-CR led to altered SIRT1 activity in liver, we analysed the lysine acetylation of two well-characterized SIRT1 targets, p53 (on K379) and FOXO1 (on K242, K245 and K262)\(^{15–17}\). We observed increased lysine acetylation of both targets in TRF-CR compared with TRF-Reg mice during the daytime, similar to hyperacetylation of these TFs in liver-specific Sirt1 knockout mice (L-Sirt1\(^{−/−}\)) (Fig. 3d and Extended Data Fig. 3e). Further, recruitment of FOXO1 to DNA was reduced during the day in TRF-CR mice, similar to the decreased genome-wide FOXO1 binding observed in L-Sirt1\(^{−/−}\) liver (Extended Data Fig. 3d)\(^{19,20}\).

Transduction of cytonuclear LbNOX in the liver during TRF-CR reduced FOXO1 and p53 acetylation during the daytime but not the night-time, consistent with daytime activation of SIRT1 by NOX during TRF-CR (Fig. 3d, right). We also note that both control mice subjected to TRF-CR and L-Sirt1 knockout mice on TRF-CR have similar body temperatures, suggesting that decreased activity of SIRT1 results in lower core body temperature under TRF-CR. Together, these findings show that hepatic NADH drives rhythmic energy conservation during TRF-CR by inhibiting SIRT1.

**Discussion**

Results presented here reveal that the circadian system participates in the response to a low-energy state associated with hypocaloric feeding through the rhythmic elevation of NADH during the daytime and inhibition of SIRT1 in liver. In nocturnal rodents provided a CR diet limited to the active period, we observed that daytime NADH elevation drives a daily drop in core body temperature in the morning, rhythmic inhibition of SIRT1, decreased BMAL1-dependent gene expression and reduced acyl-carnitine and amino acid levels. We speculate that NADH regulates energy conservation in the morning when food becomes scarce for nocturnal mice through the inhibition of SIRT1-controlled gene networks (Fig. 4).

Overexpression of the cytonuclear NADH-specific oxidase LbNOX, in combination with liver-specific ablation of Sirt1, has enabled us to dissociate hepatic redox and nutritional state, providing insight into how NADH controls SIRT1 in vivo during the daily fasting period\(^{44}\). We find that alleviation of NADH inhibition of SIRT1 during the daytime leads to genome-wide disinhibition of SIRT1-controlled RNAs during TRF-CR. In addition, liver isolated from LbNOX-transduced animals was enriched in PPAR\(\alpha\)-regulated transcripts, a major transcription factor involved in the switch to oxidative metabolism during fasting\(^{41}\). It is likely that downregulation of PPAR\(\alpha\) and BMAL1 in the morning/fasting period during TRF-CR contributes to the daily rhythms in energy conservation characterized by reduced body temperature and entry into torpor during the fasting period\(^{43}\). Indeed, downregulation of hepatic fatty acid oxidation and metabolic transcriptional reprogramming via the core circadian transcription factors BMAL1 and CLOCK, acting in concert with PPAR\(\alpha\), leads to daily rhythms in body temperature and body weight in the absence of food (Extended Data Fig. 4).
acid oxidation has recently been shown to exacerbate cold sensitivity\textsuperscript{64}. Finally, identification of the methionine pathway downstream of NADH/SIRT1/BMAL1 during TRF-CR raises the possibility that the healthful effects of both calorie and methionine restriction may be mediated through a common mechanism\textsuperscript{61–63}. Strategies to mimic intermittent NADH redox inhibition of SIRT1 in liver may recapitulate beneficial effects observed with TRF and be leveraged towards altering fuel selection in cells type involved in proliferative and inflammatory disease states.

Although sirtuins have been associated with the beneficial effects of CR across a range of organisms\textsuperscript{22,46,65} and CR promotes longevity in yeast by increasing \( \gamma \)-Sir2p activity and NAD\(^+\)/NADH\textsuperscript{4,6}, the situation is more complex in mammals, because CR exerts tissue-specific effects on NAD\(^+\) and NADH\textsuperscript{22,46,67}. Our findings using SAMDI-MS indicate that NADH inhibits SIRT1 activity at physiologically relevant concentrations\textsuperscript{22,25–26}, and our in vivo genetic analyses indicate that hepatic NADH inhibition of SIRT1 during the daytime plays an important role in energy-sparing in liver under a low-energy state. Whether redox regulation of other sirtuin isoforms (SIRT2–7) could also contribute to tissue-specific promotion of energy conservation during low-energy conditions remains to be explored. Future studies are necessary to elucidate whether other energy-sensing pathways influence the capacity of NADH to regulate the activity of SIRT1 or other sirtuin isoforms during low-energy states in vivo. Our finding that the cytonuclear form of \( LbNOX \) drives SIRT1 activity in the nucleus to affect the transcriptional, metabolic and physiological response to TRF-CR suggests that SIRT1 is sensitive to changes in cytosolic NADH and/or to metabolic conditions that cause export of mitochondrial NADH equivalents into the cytosol as occurs during gluconeogenesis\textsuperscript{68–69}. Although recent reports suggest that SIRT1 may interact with NADH in the nucleus,\textsuperscript{70} continued development of methods that can accurately report subcellular concentrations of reduced/oxidized NAD(H) in vivo will facilitate greater understanding of how redox state across the day under different nutrient conditions may regulate SIRT1 and other sirtuin isoforms\textsuperscript{71}.

**Methods**

**Animals.** All animal procedures were in accordance with guidelines of the Institutional Animal Care and Use Committee at Northwestern University. Mouse protocols approved in this study include IS00003543, IS00007712, IS00001143 and IS00006001. Mice were housed at 23–25 °C in the Center for Comparative Medicine at Northwestern University and maintained under 12:12 h light/dark cycles at 40–60% humidity with ad libitum access to regular chow and water unless otherwise indicated. C57BL/6J mice were purchased from Jackson Laboratories. SIRT1\textsuperscript{lox/lox} mice harbouring \( LoxP \) sites surrounding exon 4 of \( Sirt1 \) were from S. Imai (Washington University, St. Louis). \( Abl-Cre \) mice were purchased from Jackson Laboratories. For liver-specific \( Sirt1 \) knockout studies, \( Abl-Cre; Sirt1\textsuperscript{f/f} \) mice were used for western blotting and CHIP-Seq, whereas \( Sirt1\textsuperscript{fl/fl} \) mice retro-orbitally injected with AAV8–TBG–iCre were used for transcriptomic, metabolomic and physiological studies. For inducible liver-specific \( Bmal1 \) knockout studies, mice harbouring \( LoxP \) sites surrounding exon 8 were retro-orbitally injected with AAV8–TBG–iCre. Male mice, 4–6 months old, were used in all experiments, except for studies monitoring fasted body temperature which were performed in 4–6-month-old female mice.

**Caloric restriction studies.** Mice were housed individually in cages containing automated, programmable feeder devices (Actimetrics) as described previously\textsuperscript{10}. Mice acclimated to feeder cages for 3 days with ad libitum access to food, where the average mouse consumed 1100 mg of food each day. During dietary intervention, mice fed regular Chow received 1100 mg of food daily. Mice fed regular Chow received 1100 mg of food daily. Mice were divided into 4 groups: control, ad lib, 40% DR and 30% DR groups, with 12 mice per group. Mice in the control group were fed regular Chow daily, whereas the 40% and 30% DR groups were fed regular Chow to maintain intracellular equilibrium as described previously\textsuperscript{10}. Intraperitoneal injections were performed into mice at ZT4 (daytime) to achieve a dose of 1 kg\textsuperscript{-1}. Liver was collected after 30 min for MS analysis of metabolic metabolites as below. Percentage of each mass isomotomer that was detected for glyceroldehyde–3-phosphate/dihydroxyacetone phosphate was quantified.

**NAD\(^+\) and NADH quantification.** NAD\(^+\) was quantified as described previously\textsuperscript{72}. Briefly, cut and weighing tissue was homogenized in perchloric acid (Sigma) in a TissueLyzer (Qiagen). Following neutralization with K\textsubscript{2}CO\textsubscript{3}, supernatant was dialyzed 1:1 in the mobile phase and analysed by HPLC (Shimadzu) on a Supelco LC\(_2\), column (Sigma) with an ultraviolet–visible spectroscopy (UV–Vis) detector at 260 nm. NADH was extracted as described previously\textsuperscript{73}. Briefly, cut and weighed tissue was gently homogenized by plastic pellet in five volumes of nitrogen-sparged alkaline extraction buffer (25 mM NH\(_4\)Ac, 25 mM NaOH, 100 mM \( \gamma \)-acetoin). Tissue and protein precipitates were pellet and supernatant was passed through a 0.2 μm HPLC-grade nylon filter. Extract was analysed by HPLC as above with a UV–Vis detector set to 340 nm.

**Mass spectrometry for hydrophilic metabolites.** Hydrophilic metabolites were analysed by HPLC–MS/MS as described previously\textsuperscript{73}. Briefly, 20–100 mg of weighed tissue was homogenized in 1 ml of cold methanol in the TissueLyzer, centrifuged and the equivalent volume for 10 mg of tissue was dried by SpeedVac, reconstituted in 50 mM acetonitrile and applied to the HPLC–MS/MS analysis. Data acquisition and analysis were carried out using Xcalibur v4.1 and Tracefinder v4.1 software, respectively (Thermo Fisher Scientific). Metabolite concentrations within each sample were normalized to total ion count. Data are represented as log(fold change) relative to the control condition as indicated in the text and legends.

**Body temperature analyses.** Body temperature was monitored as described previously\textsuperscript{74}. Briefly, mice were subcutaneously implanted with wireless temperature probes (IPTT-300) in the dorsal region. After a 1-week recovery, animals were monitored from outside the cage with a handheld device (DAS-8007-1US) every 2 h for either 24 or 48 h with minimal disruption to the mice (Bio Medic Data Systems). For analyses of fasted liver-specific \( Bmal1 \) or \( Sirt1 \) knockout female mice, mice were first moved into fresh cages without food at ZT0 (lights on), and body temperature was monitored as above.

**Retro-orbital delivery of AAVs.** Retro-orbital delivery of AAVs was performed as described previously\textsuperscript{75}. Briefly, \( LoxNOS \) and control plasmids (Addgene 75285) were cloned into AAV expression vectors under the thyroid-binding globulin (TBG) promoter (to avoid inflammation associated with adenoviral-based methods), packaged into AAVs of serotype 8, purified, and concentrated by Vector Biolabs. AAV8–TBG–iCre and AAV8–TBG–green fluorescence protein (GFP) plasmids were purchased from Vector Biolabs (catalogue numbers: V81724 and V81743, respectively). Following isoformease aesthesia, mice were injected retro-orbitally with 1 × 10\(^{12} \) genome copies (GC) of the respective virus in 100 μl. For mice expressing multiple AAVs, mice were co-injected with a 100 μl mixture of both viruses to achieve 5 × 10\(^{10} \) GC of iCre/GFP AAVs and 1 × 10\(^{10} \) GC of LoxNOS null AAVs. Mice were monitored acutely for recovery after anaesthesia then given 3 days to recover from AAVs administration. Mice were then transferred to experimental chambers and allowed to acclimate to that environment for 2 weeks before the start of dietary interventions.

**RNA sequencing and analysis.** RNA was isolated from 10–20 mg of liver tissue using the Zymo Direct-Zol RNA miniPrep kit as described described\textsuperscript{76}. RNAs
with an RNA integrity number > 8 were used for library preparation, which was constructed from 230 ng of RNA with the NEBNext RNA Ultra Directional library prep kit (NEB). Seventy-five base pair single-end sequencing was performed on a NextSeq 500 sequencer with a high output Illumina flow cell (FC-404-3002) and analysed as described previously25. Briefly, sequences were aligned to the mm10 genome with STAR (v.2.5.2), assigned to Ensembl features (GRCm38.60.12) with subread:featureCounts (v.1.5.1.1), and analysed for differential expression at each time point in DESeq2 (v.1.24.0). The FDR-adjusted P-value cut-off for significance was set at 0.05. Gene ontology Kyoto Encyclopedia of Genes and Genomes (KEGG) analyses were performed with the HOMER find GO command, and principal components analyses were performed by DESeq2 on variance stabilization transformed RNA-Seq data for genes differentially expressed by diet. Known motif analysis was performed by HOMER (v.4.8.3.3) using the JASPAR database of DNA-binding motifs for regions of open chromatin previously identified by ATAC-seq, sequencing in liver of ad libitum mice at ZT4 that annotate to genes listed as in results and legends. For comparison of genomic datasets, quadrant plots were utilized in which log2(fold change) in gene expression from TRF-CR (relative to TRF-Reg) was compared with the log2(fold change) in gene expression from the given dataset using genes that are significantly differentially expressed by diet. Genes within each quadrant were counted and expressed as a fraction of the total. For comparison with outside datasets involving Bmal1−/− and Ppara−/−, a fold change cut-off of ±0.5 was applied. LNNOX and iCre expression were determined by using Bowtie2 (v.2.2.4) to align to their respective sequences and dividing by ten million reads of each library. Sirt1 knockout was confirmed by taking the ratio of reads within the exon flanked by LoxP sites (exon 4) and exon 9 of Sirt1 using subread:featureCounts set to count by exon.

ChIP sequencing and analysis. Liver was processed for ChIP-Seq as described previously16. Briefly, liver was double cross-linked, nuclei isolated with hypotonic buffer and needle lysis, and chromatin sheared by sonication (Diagenode). Chromatin was diluted 1:10 in dilution buffer (0.1% SDS, 1.1% Triton X-100, 167 mM NaCl, 1.2 mM EDTA pH 8, 1.67 mM Tris–HCl pH 8), and immunoprecipitations were performed with 15 μg of anti-BMAL1 (Millipore, catalogue number: ABE2599) or FOXO1 (Abcam, catalogue number: ab39670) antibodies, and secondary-conjugated, bovine serum albumin-blocked paramagnetic beads pulldown protein/DNA complexes. Chromatin was de-cross-linked and purified before library prep (NEBNext Ultra II library prep kit). 2 ng of each replicate was processed into libraries, which were size selected to 200–600 bp before polymerase chain reaction amplification and pooled for sequencing as in ref.16. Bowtie2 (v.2.2.4.2) was used to align sequencing data to the mm10 genome with standard parameters. Peaks were called using the HOMER (v.4.8.3) findPeaks command with the settings -style factor, -size 275, -fragLength 250. For scatter plots, the tag-density for individual peaks for each replicate as described was quantified by HOMER annotatePeaks with setting -size, given, and averaged. For quadrant plots, a fold change cut-off of ±0.5 was applied and the number of peaks within each quadrant was counted.

Solid-phase peptide synthesis. Peptide substrates were synthesized on Fmoc-ribose amide 4-methylbenzhydrylamidine resin (AnaSpec, Inc.) as described previously16. Briefly, the N-terminal fluorenylmethyloxycarbonyl was deprotected with 20% piperidine in N,N-dimethylformamide, filtered and washed with N,N-dimethylformamide,对抗meptide beads pulldown protein/DNA complexes. Chromatin was de-cross-linked and purified before library prep (NEBNext Ultra II library prep kit). 2 ng of each replicate was processed into libraries, which were size selected to 200–600 bp before polymerase chain reaction amplification and pooled for sequencing as in ref.16. Bowtie2 (v.2.2.4.2) was used to align sequencing data to the mm10 genome with standard parameters. Peaks were called using the HOMER (v.4.8.3) findPeaks command with the settings -style factor, -size 275, -fragLength 250. For scatter plots, the tag-density for individual peaks for each replicate as described was quantified by HOMER annotatePeaks with setting -size, given, and averaged. For quadrant plots, a fold change cut-off of ±0.5 was applied and the number of peaks within each quadrant was counted.

Protein expression. SIRT1 (amino acids 193–747) was produced as a recombinant protein in E. coli (strain BL21 DE3) using a pTriEx-based Protein expression vector. Cells were grown in 2XTY broth supplied with 100 μg/mL kanamycin and 50 μg/mL carbenicillin at 37 °C until the optical density reached ~0.6, then induced with isopropyl-β-D-thiogalactoside at 37 °C for 3 h. Protein extracts were subjected to SDS–PAGE, transferred to nitrocellulose membranes and probed with primary antibodies specific for acetyl-lysine targets regulated by SIRT1 (H3K9-acetyl; Abcam, catalogue number: ab88989; H4K16-acetyl; Millipore, catalogue number: 07-329; FOXO1 (K242, K245, K292)–acetyl; Santa Cruz, catalogue number: sc3744272; P33(K379)-acetyl (Cell Signaling, catalogue number: 2570)), as well as total levels of H3 (Cell Signaling, catalogue number: 9715), H4 (Cell Signaling, catalogue number: 13919), FOXO1 (Santa Cruz, catalogue number: sc11350), P33 (Cell Signaling, catalogue number: 2524), SIRT1 (Millipore, catalogue number: 07-131), FLAG (Sigma, catalogue number: M8823) and ACTIN (Cell Signaling, catalogue number: 4970) at a dilution of 1:1,000. Signal was quantified relative to total H3 or ACTIN by performing background-corrected densitometry in ImageJ (v.2.0.0). Uncropped western blot scans labelled with molecular weight markers are presented in the Source Data files.

Acyl-carnitine quantitation. Acyl-carnitine measurements were made by flow-injection MS/MS using sample preparation methods described previously29,30. Data was acquired using a Waters TQD mass spectrometer equipped with an Acquity UPLC system and controlled by MassLynx 4.1 operating system (Waters).

Immunohistochemistry. Immunohistochemistry was performed with FLAG antibody (Sigma). Briefly, livers were fresh-fixed in 4% paraformaldehyde before embedding in OCT compound (Tissue Tek) and freezing at −80 °C. Liver slices (20 μm thick) were prepared on a cryostat and stained with FLAG antibody at 1:100 dilution before bright-field colour imaging at ×4 magnification.

Public genomics datasets used in this study. RNA-Seq data from liver of ad libitum PPARα controls: GSE118787. RNA-Seq data from liver of WT and Bmal1−/− mice: GSE133899.

Quantitation and statistical analysis. Throughout the article, log2(fold change (TRF-CR)) indicates TRF-CR relative to TRF-Reg for null mice unless a different genotype is indicated; log2(fold change (LNNOX in TRF-CR)) indicates TRF-CR, LNNOX relative to TRF-CR, null; log2(fold change (LNNOX in TRF-CR, L-Strit−/−)) indicates TRF-CR, LNNOX relative to TRF-CR, L-Strit−/−, null; log2(fold change (Bmal1−/−)) indicates Bmal1−/− relative to wild-type littermates; log2(fold change (Pyaro−/−)) indicates Pyaro−/− relative to wild-type littermates. Statistical analysis was performed with unpaired two-tailed Student’s t test in Microsoft Excel (v.16.16.16) unless noted otherwise. Data are represented as mean ± s.e.m. unless noted otherwise. Biologically independent replicates in each experiment are noted in the figure legends as the ‘n’ specific to each group. Only one ‘n’ is provided if the number of independent biological replicates is the same across groups. Quadrant plots were analysed by binomial test. Box and whisker plots depict the following: line, median; box limits, first and third quartiles; whiskers, 10th and 90th percentiles; outliers, not shown. Differences were considered significant when P < 0.05 unless noted otherwise. Body temperature rhythms were analysed by two-way analysis of variance, and average core body temperature at each sampling time. The final enzyme concentrations of ~30–100 nM, with NAD+ and NADH as indicated. Then 1 μl of the acetylated peptide substrate was added at a final concentration of the peptide’s Michaelis constant (Km) to initiate the reaction. The reaction plate was incubated at 37 °C for 15–20 min (within the linear response range), then quenched with excess of the deacetylase inhibitor nicotinamide (50 mM). KmNADH and KmNAD+ were derived by fitting the initial velocity data into a nonlinear regression competitive inhibitor model in Graphpad Prism software (v.9.2.0). We derived the kinetic constants assuming the kinetics of the thiol form of substrate and the disulfide form are the same. Lineweaver–Burk transformation was determined by inverting data and conducting linear curve-fitting.
analysis of variance. Differentially expressed liver TRF-CR genes, liver metabolites and serum acyl-carnitines that require SIRT1 for the effect of LBNOX were those that demonstrated either: (1) log(log change (TRF-CR)) > 0 and log(log change (LBNOX in TRF-CR) < 0 and log(log change (LBNOX in TRF-CR, log10(1+TRF-CR)) > 0.5 × log(log change (LBNOX in TRF-CR)); or (2) log(log change (TRF-CR) < 0 and log(log change (LBNOX in TRF-CR)) > 0 and log(log change (LBNOX in TRF-CR, log10(1−TRF-CR)) < 0.5 × log(log change (LBNOX in TRF-CR)).

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
Data generated in this study are publicly available in the GEO repository (GSE151281). We also utilized publicly accessible RNA-seq data from GEO repositories GSE133989 and GSE118787; JASPAR databases are found at http://jaspar.genereg.net/search?q=collection=CORE&tax_group=vertebrates. Correspondence and requests for materials should be addressed to Joseph Bass (j-bass@northwestern.edu). Source data are provided with this paper.

Received: 11 August 2021; Accepted: 28 October 2021; Published online: 13 December 2021

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Extended Data Fig. 1 | See next page for caption.
Extended Data Fig. 1 | Circadian and metabolic profiling in TRF-CR mice. a, (left) Representative wheel-running behaviour and (middle) quantification of average daily onset of activity for TRF-Reg (n = 5) and TRF-CR (n = 5) mice throughout the duration of the 4-wk intervention (double plotted for clarity). (right) Background-corrected PER2::LUCIFERASE readings from excised suprachiasmatic nucleus (SCN) of TRF-Reg (n = 6) and TRF-CR (n = 6) mice. The grey shading indicates mean values ± SEM. Experiments were performed in 4–6 mo old male C57BL6/J mice. b, Body weight of TRF-Reg (n = 15) and TRF-CR (n = 15) mice over the duration of the 4-wk study. c, Blood glucose and serum insulin during oral glucose tolerance testing performed during the daytime (at ZT4) in TRF-Reg (n = 6) and TRF-CR (n = 6) mice. d, GAP/DHAP mass isotopomer distribution determined by mass spectrometry of liver from TRF-Reg (n = 3) and TRF-CR (n = 2) mice 30 min following i.p. administration of a 10:1 mix of U13C-lactate:U13C-pyruvate (1g/kg). e–f, (left) Relative concentration of (e) NADH and (f) NAD+ by HPLC in liver of TRF-CR compared to TRF-Reg mice during the day (ZT4) (n = 12 for TRF-Reg, n = 11 for TRF-CR) and at night (ZT16) (n = 9 for each diet) for each time point. (right) Concentration of (e) NADH and (f) NAD+ in ad lib fed wild-type mouse liver by HPLC every 6 hrs for 24 hrs (n = 7). g–h, Log2-FC in daytime liver acyl-carnitine levels in (g) TRF-CR compared to TRF-Reg mice and (h) LbNOX-compared to null-overexpressing TRF-CR mice (n = 3). Data are presented as mean values ± SEM. Statistics were performed with unpaired, two-tailed student’s t test except as otherwise noted in the figure. *p < 0.5, ***p < 0.01.
Extended Data Fig. 2 | See next page for caption.
Extended Data Fig. 2 | Daytime NADH elevation regulates genome-wide transcription of fatty acid metabolism genes through BMAL1 during TRF-CR. 

a, (left) RNA-seq reads per 10 million sequenced reads (RPM) that align to LbNOX in LbNOX- vs null-transduced TRF-Reg (n = 6 per genotype) and TRF-CR (n = 6 for null; n = 5 for LbNOX) liver. (right) Representative in situ immunohistochemistry against FLAG-LbNOX in liver of null- and LbNOX-transduced mice. Experiments were performed in 4–6 mo old male C57BL6/J mice unless otherwise noted. 

b, Heatmap depicting log2-FC in gene expression from TRF-CR and LbNOX in TRF-CR for genes within gluconeogenic and glycolytic gene ontology groups. 

c, Unbiased principal components analysis during the day (ZT4) or night (ZT16) of null- and LbNOX-expressing TRF-Reg or TRF-CR mice for genes that are differentially-expressed by TRF-CR in null mice at respective time points (DESeq2 p-adj<0.05) (n as in panel a). 

d, Log2-transformed FC in expression of select fatty acid metabolism genes during TRF-CR (blue) (n = 6), following LbNOX-overexpression during TRF-CR (white) (n = 5), and in Bmal1 knockout liver (purple) (n = 3). 

e, 24-hour fasted body temperatures in 4–6 mo old female liver-specific Bmal1fx/fx mice before and after retro-orbital administration of AAV8-TBG-iCre (n = 6) (left: ANOVA, p < 0.01; right: *p < 0.05 in paired, two-way student’s t test). 

f, Log2-transformed FC in expression of genes within the methionine pathway for conditions (n as in panel d). 

g, (top) Summary of methionine pathway and (bottom) metabolomics of effect of TRF-CR and LbNOX in TRF-CR on end-products of methionine metabolism in liver and serum performed during the daytime (ZT4) (n = 3). Data are presented as mean values ± SEM.
Extended Data Fig. 3 | See next page for caption.
Extended Data Fig. 3 | NADH inhibition of SIRT1 activity. a, Lineweaver-Burk transformation of data from SAMDI-MS. b, Relative NADH quantified by HPLC following supplementation with pyruvate or lactate compared to controls (n = 3). c, Densitometric quantification of western blots from liver of 4–6 mo old male mice in indicated genotypes at ZT4 for pS3-(K379)Ac (left) and FOXO1-(K242, K245, K262)Ac (right) relative to control TRF-Reg (from Fig. 3d) (n = 3). d, FOXO1 ChIP-seq from liver collected during the daytime comparing effect of FOXO1 binding in TRF-CR (x-axis) with the effect of hepatic SIRT1 ablation (L-Sirt1−/−) (y-axis). Each point indicates a FOXO1 peak in control liver. Peaks that have absolute log_{2}(fold change) >0.5 for both comparisons are colored blue or black and counted by quadrant (n = 2). e, Densitometric quantification of western blots as in (c) relative to null-transduced TRF-Reg mice (from Fig. 3d) (Ac-pS3: n = 4 for TRF-CR, LbNOX. n = 6 for all other conditions; Ac-FOXO1: n = 2 for TRF-CR, LbNOX. n = 3 for all other conditions). f, FOXO1 ChIP-seq from liver during the daytime comparing effect of TRF-CR (x-axis) with effect of hepatic SIRT1 ablation (L-Sirt1−/−) (y-axis) (n = 2). g, Western blotting for Ac-FOXO1 during the nighttime (ZT16) in null- and LbNOX-transduced mice on TRF-Reg and TRF-CR (n.s. non-specific) and densitometric quantification (n = 3). h, BMAL1 ChIP-seq from liver during the daytime comparing the effect of TRF-CR and L-Sirt1−/− as in panel d (n = 3). Uncropped Western blot scans labelled with molecular weight markers are presented in the Source Data Files. i, RNA-seq reads mapping to the exon of Sirt1 that is flanked by LoxP sites (Ex4) relative to exon 9 (Ex9) of Sirt1, and RNA-seq reads per 10 million sequenced reads (RPM) that align to iCre and LbNOX in null- or LbNOX-transduced TRF-CR mice co-transduced with iCre (n = 6). Null-expressing mice on TRF-CR (blue) (n = 5) are shown as reference. j, Quadrant plot comparing transcriptional responses to TRF-CR in null-transduced mice (x-axis) (n = 6) and LbNOX-expression in TRF-CR, L-Sirt1−/− mice (y-axis) (n = 6). Each point indicates a gene that is DE by TRF-CR in null-transduced mice (930 genes). Percentages of genes within each quadrant are shown. k, Average 48-hour fasting body temperature in 4–6 mo old female liver-specific Sirt1−/− mice (n = 5). Data are presented as mean values ± SEM. Statistics were performed with unpaired, two-tailed student’s t test except as otherwise noted in the figure. *p < 0.05.
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- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen’s d, Pearson’s r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

We used publicly-accessible software and standard parameters to analyze our data. For RNA-Seq, STAR (v2.5.2), subread:featureCounts (v1.5.1), DESeq2 (v1.24.0), and HOMER (v4.8.3) were used. For ChIP-Seq, Bowtie2 (v2.2.4) and HOMER (v4.8.3) were used. ClockLab software (Version 6, Actimetrics) was used to analyze wheel running activity, and Tracefinder (v4.1, Thermo Fisher Scientific) was used to analyze metabolite concentrations following HPLC-MS/MS. All other analyses were performed in Microsoft Excel (v16.16.16) and Graphpad PRISM (v9.2.0). Western blot images were quantified with ImageJ (v2.0.0).

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

Data

Policy information about availability of data

All manuscripts must include a data availability statement. This statement should provide the following information, where applicable:
- Accession codes, unique identifiers, or web links for publicly available datasets
- A list of figures that have associated raw data
- A description of any restrictions on data availability

Data and Materials Availability: Data generated in this study are publicly available in the GEO repository (GSE151281). We also utilized publicly-accessible RNA-seq data from GEO repositories GSE133989 and GSE118787. JASPAR databases are found at http://jaspar.genereg.net/search?
Field-specific reporting

Please select the one below that is the best fit for your research. If you are not sure, read the appropriate sections before making your selection.

- Life sciences
- Behavioural & social sciences
- Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see [nature.com/documents/nr-reporting-summary-flat.pdf](http://nature.com/documents/nr-reporting-summary-flat.pdf)

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | Power calculations were performed to identify the minimum number of mice for each assay that would achieve 90% power given measured differences in means and standard deviations for each assay in previous work in our lab as in (Levine DC et al, Molecular Cell, 2020). |
|-------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Data exclusions | We used pre-determined criteria to exclude outliers including IQR*1.5 rule or PCA clustering at a substantial distance away from other replicates. One replicate from the AAV8-IbNOX-transduced, TRF-CR group was excluded as a result of being identified as an outlier by the above criteria in assays measuring protein and RNA levels of IbNOX, NADH, and also in clustering of RNA-seq data. |
| Replication | All findings in mice were successfully replicated at each attempt and were replicated in at least one independent cohort containing multiple mice. In cell-based studies, results were repeated in at least 3 independent experiments collected and assayed on different days. |
| Randomization | Littermates were distributed evenly into each group. |
| Blinding | Code-names were applied to samples to blind researchers during sample processing at the bench and during data analysis. |

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

**Materials & experimental systems**

| n/a | Involved in the study |
|-----|-----------------------|
| ☐ | Antibodies |
| ☒ | Eukaryotic cell lines |
| ☐ | Palaeontology and archaeology |
| ☒ | Animals and other organisms |
| ☐ | Human research participants |
| ☒ | Clinical data |
| ☐ | Dual use research of concern |

**Methods**

| n/a | Involved in the study |
|-----|-----------------------|
| ☒ | ChIP-seq |
| ☐ | Flow cytometry |
| ☒ | MRI-based neuroimaging |

**Antibodies**

| Antibodies used | All antibodies were used at a dilution of 1:1000. H3K9-Ac (Abcam ab8898), H4K16-Ac (Millipore 07-329), FOXO1(K242,K245,K262)-Ac (Santa Cruz sc-49437), p53(K379)-Ac (Cell Signaling 2570), H3 (Cell Signaling 9715), H4 (Cell Signaling 13919), FOXO1 (Santa Cruz sc374427), p53 (Cell Signaling 2524), SIRT1 (Millipore 07-131), FLAG (Sigma M8823), ACTIN (Cell Signaling 4970), FOXO1 ChIP (Abcam ab39670), and BMAL1 ChIP (Millipore ABE2599). |
| Validation | Antibodies chosen have been widely published, contain validation statements in the vendor websites, and/or were validated by the Antibody Validation Database (Park Lab). For the FOXO1 Abcam ChIP antibody, ChIP-seq datasets were validated by confirming expected phenotypes (i.e. increased binding to known targets following prolonged fasting and in insulin receptor knockouts), by performing unbiased DNA motif analysis with HOMER, and by comparing results to a Santa Cruz FOXO1 antibody (PMID: 30532187) using peaks from our antibody as reference. |

**Eukaryotic cell lines**

| Cell line source(s) | Immortal wild type mouse embryonic fibroblasts were generated in lab as previously described (Levine DC et al, Molecular Cell, 2020). HEK293T cells were obtained from Takara (632180). |
Authentication

None of the cell lines were authenticated.

Mycoplasma contamination

MEF and HEK293 cells used in this study have tested negative for mycoplasma contamination.

Commonly misidentified lines

Our cell lines are not commonly misidentified.

## Animals and other organisms

**Policy information about studies involving animals; ARRIVE guidelines** recommended for reporting animal research

**Laboratory animals**

4-6 mo old male C57B6/J mice were used in TRF-Reg and TRF-CR mice +/- LbNox. We also utilized liver-specific Sirt1 KO mice (generated either by crossing Sirt1 fx/fx mice with Alb-Cre mice or by injecting Sirt1 fx/fx mice with AAV8-TBG-iCre) and liver-specific Bmal1 KO mice (generated by injecting Bmal1 fx/fx mice with AAV8-TBG-iCre). Male 4-6 month old male mice were used in all experiments, except for studies monitoring fasted body temperature which were performed in 4-6 month old female mice.

**Wild animals**

No wild animals were used in this study.

**Field-collected samples**

No samples were collected in the field in this study.

**Ethics oversight**

All animal procedures were in accordance with guidelines of the Institutional Animal Care and Use Committee at Northwestern University. Mouse protocols approved in this study include: IS00003543, IS00007712, IS00001143, and IS00000601. Note that full information on the approval of the study protocol must also be provided in the manuscript.

## ChIP-seq

**Data deposition**

- Confirm that both raw and final processed data have been deposited in a public database such as GEO.
- Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

**Data access links**

Data in this study is publicly available in the GEO repository (GSE151281). Correspondence and requests for materials should be addressed to Joseph Bass (j-bass@northwestern.edu).

**Files in database submission**

- LbCR4-RNA.NULL.Reg.1.fastq.gz
- LbCR4-RNA.NULL.Reg.2.fastq.gz
- LbCR4-RNA.NULL.Reg.3.fastq.gz
- LbCR4-RNA.NULL.Reg.4.fastq.gz
- LbCR4-RNA.NULL.Reg.5.fastq.gz
- LbCR4-RNA.LbNOX.Reg.1.fastq.gz
- LbCR4-RNA.LbNOX.Reg.2.fastq.gz
- LbCR4-RNA.LbNOX.Reg.3.fastq.gz
- LbCR4-RNA.LbNOX.Reg.4.fastq.gz
- LbCR4-NULL.LbNOX.Reg.5.fastq.gz
- LbCR4-NULL.LbNOX.Reg.6.fastq.gz
- LbCR4-NULL.CR.1.fastq.gz
- LbCR4-NULL.CR.2.fastq.gz
- LbCR4-NULL.CR.3.fastq.gz
- LbCR4-NULL.CR.4.fastq.gz
- LbCR4-NULL.CR.5.fastq.gz
- LbCR4-NULL.CR.6.fastq.gz
- LbCR4-NULL.CR.7.fastq.gz
- LbCR4-LbNOX.CR.1.fastq.gz
- LbCR4-LbNOX.CR.2.fastq.gz
- LbCR4-LbNOX.CR.3.fastq.gz
- LbCR4-LbNOX.CR.4.fastq.gz
- LbCR4-LbNOX.CR.5.fastq.gz
- LbCR4-FOXO1.NULL.Reg.1.fastq.gz
- LbCR4-FOXO1.NULL.Reg.2.fastq.gz
- LbCR4-FOXO1.NULL.Reg.3.fastq.gz
- LbCR4-FOXO1.NULL.Reg.4.fastq.gz
- LbCR4-FOXO1.NULL.Reg.5.fastq.gz
- LbCR4-FOXO1.NULL.Reg.6.fastq.gz
- LbCR4-FOXO1.NULL.CR.1.fastq.gz
- LbCR4-FOXO1.NULL.CR.2.fastq.gz
- LbCR4-FOXO1.NULL.CR.3.fastq.gz
- LbCR4-FOXO1.NULL.CR.4.fastq.gz
- LbCR4-FOXO1.NULL.CR.5.fastq.gz
- LbCR4-FOXO1.NULL.CR.6.fastq.gz
- S1CR4-FOXO1.CTRL.Reg.1.fastq.gz
- S1CR4-FOXO1.CTRL.Reg.2.fastq.gz
- S1CR4-FOXO1.CTRL.CR.1.fastq.gz
- S1CR4-FOXO1.CTRL.CR.2.fastq.gz
- S1CR4-FOXO1.LSKO.Reg.1.fastq.gz
- S1CR4-FOXO1.LSKO.Reg.2.fastq.gz
Genome browser session
(e.g. UCSC)

Genome browser session
no longer applicable

Methodology

Replicates

Exact replicate numbers are stated in the figure legends. Typically, separate ChIPs, library preps, and analyses were performed for each of 2-3 mice with signal averaged during plotting. In some cases the number of mice was higher. Replicates were tested for agreement by assessing deviation from the diagonal in a scatterplot.

Sequencing depth

For ChIP-Seq, we performed 75 base pair, single-end sequencing to generate ~15-25M reads per sample, of which, ~70-80% aligned to the mm10 genome exactly 1 time.

Antibodies

Immunoprecipitations were performed in 1/4 liver with 15 μg of anti-BMAL1 (Millipore ABE2599) or FOXO1 (Abcam ab39670).

Peak calling parameters

For each ChIP, peaks were called using the HOMER findPeaks command with settings -style factor, -size 275, -fragLength 250. We used the HOMER defaults enrichment parameters which include a 4-fold enrichment over background (input), a 4-fold enrichment over local tags, and an FDR cutoff of 1%. Pooled inputs for each condition were used as background. Peaks over input were identified in each replicate, then merged. With these parameters, we identified ~550 FOXO1 peaks, similar to what has been reported previously (PMID: 30532187) and ~35,000 BMAL1 peaks, similar to what has been reported previously (PMID: 32369735).

Data quality

High quality data was confirmed by visually analyzing bigwig files in UCSC, performing motif-finding analysis with HOMER, confirming previously published phenotypes and binding locations, using Pippin Prep library size selection, Agilent Bioanalyzer, and standard Illumina QC parameters. We used the HOMER defaults that include a 4-fold enrichment over background (input), a 4-fold enrichment over local tags, and an FDR cutoff of 1%. With these parameters, we identified ~550 FOXO1 peaks, similar to what has been reported previously (PMID: 30532187), and ~35,000 BMAL1 peaks, similar to what has been reported previously (PMID: 32369735).
Bowtie2 (v2.2.4) was used to align sequencing data to the mm10 genome with standard parameters. For each ChIP, peaks were called using the HOMER findPeaks command with settings -style factor, -size 275, -fragLength 250. For scatter plots, tag-density for individual peaks for each replicate as described was quantified by HOMER annotatePeaks with setting -size given, and averaged. Known motif analysis was performed by HOMER using the JASPAR database of DNA binding motifs for peaks identified in results and legends. Gene ontology analysis was similarly performed with HOMER.