Transcriptome-wide analysis of PGC-1α-binding RNAs identifies genes linked to glucagon metabolic action

Clint D. J. Tavares,a,b,1 Stefan Aigner,c,d,1 Kfir Sharabi,a,b,2 Shashank Sathe,c,d, Beste Mutlu,a,b,c, Gene W. Yeo,c,d,e,3 and Pere Puigserver,a,b,3

aDepartment of Cancer Biology, Dana-Farber Cancer Institute, Boston, MA 02215; bDepartment of Cell Biology, Harvard Medical School, Boston, MA 02215; cDepartment of Cellular and Molecular Medicine, University of California San Diego, La Jolla, CA 92093; dStem Cell Program, University of California San Diego, La Jolla, CA 92093; and eInstitute for Genomic Medicine, University of California San Diego, La Jolla, CA 92093

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The peroxisome proliferator-activated receptor gamma coactivator 1-alpha (PGC-1α) is a transcriptional coactivator that controls expression of metabolic/energetic genes, programming cellular responses to nutrient and environmental adaptations such as fasting, cold, or exercise. Unlike other coactivators, PGC-1α contains protein domains involved in RNA regulation such as serine/arginine (SR) and RNA recognition motifs (RRMs). However, the RNA targets of PGC-1α and how they pertain to metabolism are unknown. To address this, we performed enhanced ultraviolet (UV) cross-linking and immunoprecipitation followed by sequencing (eCLIP-seq) in primary hepatocytes induced with glucagon. A large fraction of RNAs bound to PGC-1α were intronic sequences of genes involved in transcriptional, signaling, or metabolic function linked to glucagon and fasting responses, but were not the canonical direct transcriptional PGC-1α targets such as OXPHOS or gluconeogenic genes. Among the top-scoring RNA sequences bound to PGC-1α were Foxo1, Camk1β, Per1, Klf15, Ptn4, Cluh, Trpc5, Gfra1, and Slc25a25. PGC-1α depletion decreased a fraction of these glucagon-induced messenger RNA (mRNA) transcript levels. Importantly, knockdown of several of these genes affected glucagon-dependent glucose production, a PGC-1α-regulated metabolic pathway. These studies show that PGC-1α binds to intronic RNA sequences, some of them controlling transcript levels associated with glucagon action.

PGC-1α | RNA binding | glucagon | liver | mitochondria

The transcriptional peroxisome proliferator-activated receptor gamma coactivator 1-alpha (PGC-1α) is a canonical regulatory component of physiological processes such as cold, fasting, and exercise (1–4). PGC-1α is itself regulated at multiple levels including transcription, translation, and posttranslational (5–9). As a transcriptional coactivator, PGC-1α increases expression of genes associated with energy metabolism and mitochondrial biogenesis through binding to transcription factors (10–12). Once bound to a transcription factor it engages additional chromatin-remodeling proteins and the basal transcriptional initiation machinery to increase expression of targeted genes (13, 14). Most of the biological functions of PGC-1α have been focused on recruitment to promoters and enhancers through physical interaction with transcription factors (15). For example, PGC-1α binds to ERRα, NRFs, and YY1 to activate transcription of a large number of nuclear genes encoding for mitochondrial proteins (10, 11, 16, 17), or to HNF4α, FOXO1, and GR to augment transcription of gluconeogenic genes (2, 18–20). Interestingly, PGC-1α is one of the few transcriptional coactivators that contains serine/arginine (SR) and RNA recognition motif (RRM) domains at its C terminus, similar to splicing factors or other RNA-binding proteins (13). Some studies have reported that PGC-1α binds to components of the initial elongation machinery (21, 22). In addition, the RNA methyltransferase NSUN7 has been shown to promote PGC-1α-mediated transcription and corresponds to enrichment of a specific set of enhancer-associated transcripts (23). The C terminus of PGC-1α also binds to cap-binding protein 80 (CBP80) and both proteins appear to associate with the 5′ cap of target transcripts of promyogenic genes (24). However, none of these studies have broadly determined what type of RNAs are bound to endogenous PGC-1α in a metabolic or energetic process.

Glucagon action is a central fasting and diabetic response that controls metabolism and energy balance (25–30). Glucagon uses the ancient and canonical cyclic adenosine monophosphate pathway to control part of the fasting metabolic action response, including hepatic glucose production. Glucagon regulatory metabolic function occurs at different levels, including direct metabolic enzyme activity and fluxes and transcriptional proteins such as CREB-dependent complex assembly with the CBP/p300 and TORC2 coactivators (2, 3, 31–34). One of the CREB targets is PGC-1α that maintains gluconeogenic and fatty acid oxidation gene expression (3, 35, 36). Although PGC-1α has been shown to mediate part of the glucagon and fasting response (37–40), the complete mechanisms and targets, in particular the binding to RNAs, are not entirely understood.

Here, we have performed a transcriptome-wide analysis of PGC-1α target RNAs in glucagon-treated primary hepatocytes. Immunoprecipitated glucagon-induced endogenous PGC-1α was

Significance

Glucagon action in liver is a central response to fasting and type 2 diabetes. Glucagon action has been delineated through regulatory mechanisms involving signaling, transcription factor/coactivator-based gluconeogenic gene expression, and metabolic enzyme activity. Understanding the molecular mechanisms whereby glucagon controls energy metabolism will define new strategies and potential therapies to treat metabolic diseases. Here, we have identified a regulatory mechanism whereby PGC-1α, a known transcriptional regulator of glucagon action, binds RNAs linked to glucose energy metabolism. PGC-1α represents a class of RNA-binding proteins that act as a transcriptional coactivator through transcription factor binding, but also binds to RNA sequences to control specific mRNA transcripts encoding for metabolic and bioenergetic genes.

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PNAS Latest Articles | 1 of 10

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found in complexes that contain specific RNAs. A large fraction of these RNAs were mapped to intronic sequences of transcripts. We carried out specific analysis of a highly glucagon-induced and PGC-1α-bound RNA, the mitochondrial adenosine triphosphate (ATP) transporter SLC25A25, that is necessary for glucagon-dependent glucose production and mitochondrial energetics. These studies identify a mechanism and specific targets whereby the transcriptional coactivator PGC-1α controls glucagon action in hepatocytes.

Results
PGC-1α Binds to RNAs and Forms Protein–RNA Complexes. PGC-1α functions as a transcriptional coactivator and contains SR and RRM domains at the C terminus that are predicted to interact with RNA (Fig. 1A). To evaluate if endogenous PGC-1α indeed binds RNA, we used primary hepatocytes treated with vehicle (phosphate-buffered saline; PBS) or glucagon for 3 h (Fig. 1B). Cells were subjected to ultraviolet (UV) irradiation (254 nm at 400 mJ/cm²) to cross-link protein to nucleic acids, followed by immunoprecipitation with a validated commercial antibody specific to PGC-1α. Cross-linked samples were analyzed by Western blot showing that PGC-1α is strongly induced by glucagon, and it appeared in different molecular mass bands, both in whole-cell lysates as well as immunoprecipitates (Fig. 1C and SI Appendix, Fig. S1A). The higher molecular mass PGC-1α bands, greater than 130 kDa, were not detected in non–cross-linked samples. We arbitrarily grouped the different PGC-1α molecular mass bands, from the input (whole-cell lysate) and immunoprecipitates, into upper and lower regions (indicated by the red boxes in Fig. 1C and SI Appendix, Fig. S1A), in both the glucagon-stimulated and unstimulated hepatocyte samples. Complementary DNA (cDNA) libraries were generated from the respective RNAs cross-linked to PGC-1α in accordance with the published eCLIP (enhanced UV cross-linking and immunoprecipitation) protocol (41). Orange G staining indicates the generation of suitable sequencing libraries from the different samples for sequence analysis (Fig. 1D and SI Appendix, Fig. S1B). These experiments indicate that PGC-1α interacts with RNAs. We generated biologically duplicate libraries that were sequenced to an average depth of 13 million reads each (Fig. 2A) (42). In parallel, we also prepared and sequenced paired size-matched input control libraries (30 million reads each). Reads were mapped to the reference mouse genome and irreproducible discovery rate [IDR; $P \leq 0.001$; log$_2$(fold change) $\geq 3$] analysis (41) was performed to identify reproducible and enriched binding sites within each condition (lower vs. upper band; vehicle vs. glucagon treatment). IDR analysis identified 2,206 and 3,692 reproducible and enriched PGC-1α binding sites within the control samples for the lower and upper regions, respectively. The IDR peaks were distributed among 659 and 986 genes within the lower and upper region samples, respectively. Similarly, 413 and 2,167 sites were identified to be distributed among 118 and 716 genes within the glucagon treatment samples for the lower and upper regions.
Fig. 2. Profiling of genic regions bound by PGC-1α. (A) Scatterplots showing the correlation of read density, expressed as log2-transformed reads per million [log2(RPM)], of PGC-1α-bound transcripts between the replicates. Each data point represents a bound transcript. $R^2$, Spearman correlation coefficient. (B) The genic regions of the RNA sequences found to bind PGC-1α. (C) Glucagon stimulation in hepatocytes induces binding of specific RNAs to PGC-1α in both upper and lower complexes. (D) Boxplot of binding peak distribution across genic regions for 103 and 120 RBPs from HepG2 (blue) and K562 cells (orange), respectively, compared with PGC-1α (green). Boxes and whiskers indicate quartiles. Outliers are represented by diamonds. (E) Volcano plot of genes that are bound to PGC-1α only in response to glucagon stimulation in lower and upper PGC-1α complexes.
A Fraction of Glucagon-Dependent PGC-1α-Binding RNAs Encode for Proteins That Control Hepatic Glucose Production. We and others have previously shown that PGC-1α controls hepatic glucose production in the fasting and glucagon response through different transcription-dependent mechanisms, including regulation of gluconeogenic gene expression and tricarboxylic acid (TCA) cycle fluxes (2, 3, 31). We therefore tested whether these glucagon-induced RNA transcripts that interact with PGC-1α can control hepatic glucose production. As predicted, previously identified genes involved in this pathway, such as Camek15 and Klf15, control glucose output (47, 48), although these assays demonstrate a substrate-dependent specificity (SI Appendix, Fig. S5 C–E). Other genes, such as Gfi1, that are mildly induced by glucagon, when depleted, increased glucose production only when glycerol was used as a substrate (Fig. 3D). Importantly, depletion of Slc25α25, a mitochondrial nucleotide transporter (49), strongly suppressed glucose production, particularly upon glucagon treatment, independent of the gluconeogenic substrate usage (Fig. 3D). These experiments indicate that PGC-1α, in addition to directly increasing gluconeogenic enzyme gene expression transcription such as Pck1 or G6pc, binds to RNA transcripts that regulate glucagon-dependent hepatic glucose production.

Glucagon-Dependent PGC-1α Binding to Slc25α25 Distal Intronic and Isoform-Specific Near Exon 1 RNA Sequences Increases mRNA and Protein Expression. Based on the results described above that Slc25α25 gene expression is highly increased by glucagon, Slc25α25 transcript RNA binds to PGC-1α in a glucagon-dependent manner, and Slc25α25 is required for the glucagon action on hepatic glucose production, we decided to investigate the PGC-1α binding to Slc25α25 RNA and how Slc25α25 controls this metabolic pathway. In glucagon-treated hepatocytes, PGC-1α binds to distal intronic regions common to four out of five Slc25α25 isoforms (Fig. 4A). Interestingly, the isoforms of Slc25α25 differ in their exon 1 sequences (Fig. 4A), which encode different numbers of calcium-binding EF hands at the N terminus (49), which potentially provides differential calcium sensitivity to the specific isoforms. Importantly, there was a specific binding of PGC-1α near the exon 1 RNA sequence that is closest to the third Slc25α25 isoform (TV3) that was entirely dependent on glucagon stimulation (Fig. 4A). Interestingly, among the five Slc25α25 isoforms, glucagon specifically promotes the greatest induction of TV3 (Fig. 4B), and depletion of PGC-1α selectively decreased induction of this isoform compared with TV1 and TV2 (Fig. 4C), which translates into diminished Slc25α25 mRNA and protein amounts (Fig. 4D–F). These results show that PGC-1α controls isoform-specific Slc25α25 gene expression that coincides with specific binding to distal intronic and near exon 1 RNA sequences.

SLC25A25 Regulates ATP and GTP Levels and Controls Hepatic Glucose Production. SLC25A25 is a mitochondrial nucleotide transporter that has been shown to transport ATP (49). In order to determine whether SLC25A25 could affect ATP levels in primary hepatocytes, we performed metabolite profile analysis using primary hepatocytes. The results suggest that SLC25A25 depletion altered ATP levels, such as fructose 1,6-phosphate, dihydroxy-acetone-phosphate, and α-glyceroldehyde-3-phosphate, which were also decreased (Fig. 5A). In addition, certain TCA metabolites, such as citrate and malate, were also affected by SLC25A25 depletion (Fig. 5B), indicating that SLC25A25 controls part of the hepatic gluconeogenic metabolite changes in response to glucagon. The decreases in ATP and guanosine triphosphate (GTP) (Fig. 5C) are consistent with a decrease in hepatic glucose production when SLC25A25 is depleted (Fig. 3D). Interestingly, the magnitude of glucose production suppression was higher when the isoform TV3 was depleted.
compared with TV1 (Fig. 4G). We next evaluated whether these changes in ATP and GTP correlated with altered respiratory activity. Oxygen consumption rates (OCRs) were measured in primary hepatocytes using the Seahorse instrument (Fig. 5D). Glucagon treatment caused an increase in OCR; however, SLC25A25 knockdown significantly blunted this increase, indicating that the activity of this transporter is necessary for the glucagon-dependent oxygen consumption increases. These results suggest that the PGC-1α RNA-binding target SLC25A25 controls part of the glucagon metabolic action, including increased energetics that are necessary to maintain hepatic glucose production.

**Discussion**

The PGC-1 family agglutinates a series of proteins that function as transcriptional coactivators that respond to nutrient and environmental stresses such as cold, exercise, and fasting (15, 50). All members of this family have an N-terminal transcriptional activation domain and motifs that bind to transcription factors, for example, the LXXLL motifs that bind to the ligand-binding domain of hormone nuclear receptors (1). The canonical mechanisms whereby these proteins activate gene expression are through transcription factor-based recruitment to promoters and engagement of the basal transcriptional machinery (13, 14, 51). Some members including PGC-1α contain a C terminus with SR and RRM domains involved in RNA binding, suggesting that additional mechanisms beyond promoter and transcription factor recruitment are involved (52). In fact, previous studies have implicated some of these mechanisms through binding to transcriptional elongation factors, interaction with the RNA methyltransferase NSUN7 and enhancer-associated transcripts, and CBP80 that binds to 5’ cap transcripts (21–24). Although these are mechanisms that might provide regulation of PGC-1α target genes, the PGC-1α–bound RNAs in a particular biological response were unknown. In this manuscript, we report a transcriptome-wide analysis of PGC-1α–binding RNAs identified in response to glucagon.
Fig. 4. Glucagon-dependent PGC-1α binding to Slc25a25 distal intronic and isoform-specific near exon 1 RNA sequences increases mRNA and protein expression. (A) The Slc25a25 gene gives rise to five mRNA transcripts with three different first exons (TV1 to TV3). The specific binding peak to PGC-1α, close to exon 1 of TV3, is indicated with an arrow. (B) Slc25a25 TV3 is the primary variant that is being regulated by glucagon. Data are normalized to the level of induction of total Slc25a25. (C and D) Depletion of Ppargc1a specifically suppresses the expression of Slc25a25 TV3. (E) Western blots showing the effect of siPpargc1a on total PGC-1α and SLC25A25 protein levels. (F) Western blots following mitochondrial isolation show that the protein levels of SLC25A25 are reduced in the mitochondria following siPpargc1a. (G) Depletion of Slc25a25 TV3 using specific siRNA oligos has the strongest effect on suppressing glucagon-induced glucose production. *P < 0.05, **P < 0.01, ***P < 0.001. Error bars represent the standard error of the mean.
action, a dominant fasting axis, in hepatocytes. A large fraction of these bound regions were intronic sequences, but also included exonic and 5' splice site regions, suggesting different regulatory mechanisms. This RNA binding site location pattern seems to be unique among the RNA-binding proteins analyzed in the ENCODE project (43). Based on the location pattern of PGC-1α–bound RNAs, it seems difficult to generalize a common mechanism of action integrating the PGC-1α binding with these RNAs and specific regulation. In addition, we cannot conclusively determine at this point whether the RRM domain is required for the PGC-1α–RNA binding or rather this binding is mediated through other domains of PGC-1α or other RNA-binding proteins associated with the PGC-1α complex. Thus, deciphering general principles and additional specific mechanisms underlying PGC-1α binding to RNAs will require further in-depth studies. However, for some RNA sequences, such as the first exons in Slc25a25, PGC-1α binding might provide increases in mRNAs for a specific isoform. This regulation brings an interesting concept based on whether RNA sequences located in proximal regions of gene transcription initiation can provide specific recruitment of transcriptional coactivators to define isoform specificity and mRNA transcript expression levels. In this way, the presence of RNA sequences might function, like transcription factors, as specific recruitment of activators to increase gene expression. It is also conceivable that this mechanism together with transcription factor binding could further accelerate mRNA transcript expression. In fact, this might be the case for the glucagon-inducible gene Slc25a25 that encodes for different N-terminal isoforms, and PGC-1α precisely binds near an exon 1 sequence of an isoform with specific metabolic functions. These studies also indicate that PGC-1α controls genes through different mechanisms of action providing additional points to regulate a metabolic and energetic response. Notably, PGC-1α–binding RNAs identified were not transcripts associated with the known canonical PGC-1α targets that include nuclear genes encoding for proteins linked to mitochondrial biogenesis, OXPHOS, the TCA cycle, fatty acid oxidation, or

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Fig. 5. Slc25a25 regulates ATP and GTP levels and bioenergetics in hepatocytes. (A and B) Metabolomics analysis of primary hepatocytes following siSlc25a25. The relative levels of glycolytic/glucogenic and TCA cycle intermediates are shown. (C) siSlc25a25 reduces the levels of ATP and GTP in primary hepatocytes. (D) Basal respiration of primary hepatocytes following siSlc25a25 and glucagon stimulation. *P < 0.05, **P < 0.01, ***P < 0.001. Error bars represent the standard error of the mean.
gluconeogenesis, but some of them were linked to the glucagon and fasting response. These studies also have implications for how glucagon controls energy metabolism in hepatocytes. Glucagon-induced PGC-1α binds to a series of RNAs that are known to control glucose production, indicating that this hormone uses PGC-1α as an RNA-binding protein to increase specific gene expression. This regulation confers additional mechanisms for dynamic, temporal and spatial, control of glucagon action. PGC-1α is part of a second phase of fasting control after CREB/TORC2 and CRBP/p300 activation, reinforcing the fasting response and also providing attenuating feedback mechanisms (20, 34, 38, 39). In addition to the known transcriptional targets, including HNF4α-driven PEPCK and G6Pase (2), PGC-1α uses other modes of action binding RNAs that control the gluconeogenic process, including FOXO1, CAMK1δ, KLF15, and SLC25A25. These multiple mechanisms might function to ensure increased glucose production, a process that is required for survival during fasting. Interestingly, PGC-1α binding to a specific SLC25A25 isoform, a calcium-regulated mitochondrial transporter (49), controls its expression, providing a metabolic and energetic regulation during glucagon action. We propose that glucagon is necessary to initiate transcription of Slc25a25 and that PGC-1α, by binding the RNA of a specific transcript, provides selectivity toward expression of this transcript. The expression of a specific SLC25A25 with different EF-hand calcium binding sites at the N terminus might define the transporter kinetics to control ATP transport between the mitochondria and cytosol in response to calcium fluctuations. Remarkably, this transporter is necessary to maintain cellular ATP and GTP levels, suggesting that it controls energetic mechanisms. In fact, we show that glucagon-induced respiration is defective with SLC25A25 depletion, suggesting that the ATP demand for gluconeogenesis cannot be appropriately met in the absence of SLC25A25, thus compromising the gluconeogenic pathway from different substrates. What points of gluconeogenesis are specifically altered are unclear, but the metabolite analysis suggested that different sites of gluconeogenic substrate entry are altered. A likely point of control might be the mitochondrial pyruvate carboxylase that depends on ATP levels for enzymatic activity (53, 54). In addition, previous studies have suggested that glucagon increases calcium that promotes hepatic glucose production through the inositol-1,4,5-trisphosphate receptor (55) and CAMK2 (56). SLC25A25 is likely to be part of this regulation through direct calcium binding to the EF hands located at the N terminus of the protein. Interestingly, another strong RNA hit was Trpc5, a calcium channel that might provide calcium fluxes to control SLC25A25 during glucagon action. Future studies will define this precise mechanistic regulation in the context of fasting and how it contributes to glucose and energy homeostasis in type 2 diabetes.

Methods

Primary Hepatocyte Cultures. Primary hepatocytes from 7- to 10-wk-old male C57BL/6 mice were isolated by perfusion with liver digest medium (Life Technologies; 17703-034), followed by 70-μm mesh filtration. Percoll (Sigma; P7828) gradient centrifugation was then used to separate hepatocytes from debris and other cell types. Isolated hepatocytes were seeded (4×10^6 cells per well, 6-well plates; 2×10^5 cells per well, 12-well plates; 7.2×10^4 cells, 15-cm dish) in plating medium ( Dulbecco's modified Eagle's medium [DMEM] supplemented with 10% fetal bovine serum, 2 mM sodium pyruvate, 1 μM dexamethasone, 100 nM insulin, and 1% penicillin/streptomycin). Three hours postseeding, the medium was changed to maintenance medium (DMEM supplemented with 0.2% bovine serum albumin [BSA], 2 mM sodium pyruvate, 0.1 μM dexamethasone, 1 μM insulin, and 1% penicillin/streptomycin). Cells were cultured at 37 °C in a humidified incubator containing 5% CO2 and medium was replaced daily with fresh maintenance medium.

siRNA Transfection. siRNA oligonucleotides against mouse genes were purchased from Origene Technologies (Ppargc1a, SR427524; Slc25a25, SR412336; Gfra1, SR415185; Klf15, SR413764; Camk1δ, SR412965). Hepatocytes were transfected with a 1:1 mixture of three unique 27-mer siRNA duplexes (20 nM each, with a final experimental concentration of 10 nM), using Lipofectamine RNAiMAX (Invitrogen) by reverse transfection followed with immediate seeding of hepatocytes, according to the manufacturer's protocol and as previously described (39). The following morning, the medium was replaced with maintenance medium, accompanied by another round of forward transfection, as described earlier. After 6 h, the medium was replaced again.

Treatment of Cells with Stimuli. For treatment of primary hepatocytes with glucagon, cells were incubated overnight in starvation medium (DMEM supplemented with 0.2% BSA, 2 mM sodium pyruvate, and 1% penicillin/streptomycin). The following morning, cells were stimulated with 200 nM glucagon for 3 h (unless indicated otherwise) in starvation medium.

Single-End Enhanced Cross-Linking and Immunoprecipitation. Primary hepatocytes were cultured in 15-cm dishes as described above. Cells were stimulated with or without glucagon for 3 h (three plates for each condition, done in experimental duplicates), following which they were washed twice with ice-cold PBS (pH 7.4). Cells were subjected to UV cross-linking (254 nm at 400 mJ/cm2) to generate protein–nucleic acid cross-links as described (41).

Single-end enhanced cross-linking and immunoprecipitation (seCLIP) was performed as described (41). Briefly, lysates were generated, sonicated, and then sheared to 400 nM RNA. Two percent of each sample was stored for preparation of a parallel size-matched input (SMInput) library. The remaining lysates were immunoprecipitated. Bound RNA fragments in the immunoprecipitates (IPs) were dephosphorylated and 3′-end-ligated to an RNA adapter. Protein–RNA complexes from SMInputs and IPs were run on a sodium dodecyl sulfate (SDS) polyacrylamide gel and transferred to nitrocellulose membrane. Membrane regions were excised and RNA was released from the complexes with proteinase K. SMInput samples were dephosphorylated and 3′-end-ligated to an RNA adapter. All RNA samples (IPs and SMInputs) were reverse-transcribed with AffinityScript (Agilent). cDNAs were 5′-end-ligated to a DNA adapter. cDNA yields were quantified by qPCR and 100 to 500 fmol of libraries generated with Q5 PCR Master Mix (New England Biolabs). Libraries were sequenced on an Illumina HiSeq 4000 platform.

Mapping reads were processed as described (41). Briefly, reads were adapter-trimmed using Cutadapt (v1.14) and mapped to human-specific repetitive elements from RepBase (v18.05) by STAR (v2.4.0i) (57). Repeat-mapping reads were removed, and remaining reads were mapped to the mouse genome assembly (mm10) with STAR. PCR duplicate reads were removed using the unique molecular identifier sequences in the 5′ adapter and remaining reads were retained as “usable reads.” Peaks were called on the usable reads by CLIPper (58) and assigned to gene regions annotated in GENCODE (mm10) with the following descending priority order: coding sequence (CDS), 5′ untranslated region (UTR), 3′ UTR, proximal intron, and distal intron. Proximal intron regions are defined as extending up to 500 bp from an exon–intron junction. Each peak was normalized to the SMInput by calculating the fraction of the number of usable reads from immunoprecipitation to that of the usable reads from the SMInput. Peaks were deemed significant if 8-fold enrichment and P<0.05 by exact test (or Fisher's exact test if the observed or expected read number in the eCLIP or SMInput was below 5). Sequencing and mapping statistics are described in Datasets S1 and S2. Code is available on GitHub (https://github.com/YeoLab/clip).

Cell Lysis and Western Blotting.

Cell lysis. Cells were harvested twice in ice-cold PBS (pH 7.4) (Life Technologies) and lysed in ice-cold buffer. For whole-cell extracts, cells were lysed in buffer B (1× PBS, pH 7.4, 0.5% sodium deoxycholate [wt/vol], 0.1% SDS [wt/vol], 1 mM ethylenediaminetetraacetate [EDTA], 1 mM dithiothreitol [DTT], 1% IGE PAL [vol/vol], 5 mM NaF, 5 mM β-glycerophosphate, 5 mM sodium butyrate, and 20 mM nicotinamide), supplemented with complete EDTA-free Protease Inhibitor Mixture (Roche Diagnostics). To isolate nuclei, cells were harvested in buffer C (10 mM Heps-KOH, pH 7.9, 10 mM KCl, 1.5 mM MgCl2, 0.5 mM DTT, 0.25% IGE PAL [vol/vol], 5 mM NaF, 5 mM β-glycerophosphate, 5 mM sodium butyrate, and 20 mM nicotinamide), supplemented with protease inhibitor mixture. Cytoplasmic fractions were separated, and nuclear pellets were lysed in buffer D (20 mM Heps-KOH, pH 7.9, 125 mM NaCl, 1 mM EDTA, 1 mM DTT, 1% IGE PAL [vol/vol], 10% glycerol [vol/vol], 5 mM NaF, 5 mM β-glycerophosphate, 5 mM sodium butyrate, and 20 mM nicotinamide), supplemented with protease inhibitor mixture.

Western blot analysis. Protein samples were resolved by SDS/polyacrylamide gel electrophoresis and then transfected to polyvinylidene fluoride membranes (EMD Millipore). Membranes were blocked with either 5% BSA or nonfat dry
Glucose Production Assays. Primary hepatocytes were incubated overnight in starvation medium (DMEM supplemented with 0.2% BSA, 2 mM sodium pyruvate, and 1% penicillin/streptomycin). Hepatocytes were then washed twice in warm PBS (pH 7.4). Glucose secretion by primary hepatocytes was measured by incubating the cells for 6 h in glucose-free medium (phenol-red/glucose-free DMEM, 0.2% BSA, and 2 mM sodium pyruvate and ±20 mM sodium lactate or ±10 mM glycerol), with or without concomitant stimulation with glucagon (200 nM). Medium aliquots were collected, and glucose levels were quantified using an enzymatic-based glucose assay, according to the manufacturer's protocol (Glucose/Glucose Oxidase Assay Kit; Sigma-Aldrich). The glucose concentration was calculated based on a standard curve.

Measurement of Oxygen Consumption. Oxygen consumption of primary hepatocytes was measured using a Seahorse instrument. Primary hepatocytes were seeded on an XFe24 24-well Seahorse plate and reverse transfection was performed before the assay by seeding the cells with plating media (15,000 cells per well). The next morning, medium was changed to maintenance medium and, in the afternoon, medium was changed to low-glucose maintenance medium (2.5 mM glucose). Following overnight incubation with low-glucose maintenance medium, cells were treated with 3 h with glucagon (200 nM) in no-glucose starvation medium. After 3 h, cells were washed and medium was changed to Seahorse medium (DMEM, 0.2% BSA, 2.5 mM glucose, 2 mM pyruvate, and 150 μM oleic acid) with and without glucagon. Plates were equilibrated for 45 min in a 37 °C incubator without CO2 and oxygen consumption rates were measured using the Seahorse bioanalyzer.

Metabolomics. For metabolomics analysis, primary hepatocytes were treated in a similar manner to glucose production assays using glucose-free medium containing pyruvate/lactate for the last 4 h. Following a 4-h incubation with glucagon, medium was aspirated, cells were washed with ice-cold PBS, and 800 μL of 80% methanol solution was immediately added to each well. After a 15-min incubation at ~80 °C, cells were scraped and debris was cleared (10 min, 9,000 × g). Supernatant was transferred to a new tube and the pellet was resuspended in an additional 100 μL of 80% methanol solution. Resuspended pellets were centrifuged again (5 min, 9,000 × g) and the supernatant was combined with the previous one. The combined supernatant was dried overnight using a SpeedVac and dried pellets were resuspended in 20 μL H2O before being subjected to metabolomics profiling using the AB/SCIEX 5500 QTRAP triple quadrupole instrument.

Statistics. All data are presented as means ± SEM. One-way ANOVA tests and t tests were conducted, along with corresponding posttests, as indicated; P < 0.05 was considered significant. *P < 0.05, **P < 0.01, ***P < 0.001.

Data Availability. All supporting data have been published in the National Center for Biotechnology Information Gene Expression Omnibus (https://www.ncbi.nlm.nih.gov/geo/) under accession no. GSE152303. The RBP binding data of HepG2 and K562 cells was retrieved from www.encodeproject.org, accession code ENCSR456FVU. The code for reading and mapping peak calling is available on GitHub, https://github.com/ YeoLab/clip.

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Tavares et al.


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