Deciphering the Regulatory Logic of an Ancient, Ultraconserved Nuclear Receptor Enhancer Module

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Cooperative, synergistic gene regulation by nuclear hormone receptors can increase sensitivity and amplify cellular responses to hormones. We investigated thyroid hormone (TH) and glucocorticoid (GC) synergy on the Krüppel-like factor 9 (Klf9) gene, which codes for a zinc finger transcription factor involved in development and homeostasis of diverse tissues. We identified regions of the Xenopus and mouse Klf9 genes 5–6 kb upstream of the transcription start sites that supported synergistic transactivation by TH plus GC. Within these regions, we found an orthologous sequence of approximately 180 bp that is highly conserved among tetrapods, but absent in other chordates, and possesses chromatin marks characteristic of an enhancer element. The Xenopus and mouse approximately 180-bp DNA element conferred synergistic transactivation by hormones in transient transfection assays, so we designate this the Klf9 synergy module (KSM). We identified binding sites within the mouse KSM for TH receptor, GC receptor, and nuclear factor kB. TH strongly increased recruitment of liganded GC receptor and serine 5 phosphorylated (initiating) RNA polymerase II to chromatin at the KSM, suggesting a mechanism for transcriptional synergy. The KSM is transcribed to generate long noncoding RNAs, which are also synergistically induced by combined hormone treatment, and the KSM interacts with the Klf9 promoter and a far upstream region through chromosomal looping. Our findings support that the KSM plays a central role in hormone regulation of vertebrate Klf9 genes, it evolved in the tetrapod lineage, and has been maintained by strong stabilizing selection. (Molecular Endocrinology 29: 856–872, 2015)

Abbreviations: acH3, acetylated H3; BW, body weight; 3C, chromosomal conformation capture; CHIA-PET, chromosome interaction analysis by paired end tag sequencing; ChIP, chromatin immunoprecipitation; CHX, cycloheximide; CORT, corticosterone; DMISO, dimethylsulfoxide; DNAseI, deoxyribonuclease I; dnTR, dominant negative TR; DR +4, direct repeat with 4-base spacer; eRNA, enhancer RNA; ET(OH), ethanol; FBS, fetal bovine serum; GC, glucocorticoid; GR, GC receptor; GRE/MRE, glucocorticoid/mineralocorticoid response element; H3, histone 3; hRNA, heteronuclear RNA; HS, half site; Klf9, Krüppel-like factor 9; KSM, Klf9 synergy module; IncRNA, long noncoding RNA; LPS, lipopolysaccharide; NFkB, nuclear factor kB; NR, nuclear hormone receptor; NR5, normal rabbit serum; p50, NFkB p50 subunit; PMD, postnatal day; pol II, polymerase II; RE, restriction enzyme; rpL8, ribosomal protein L8; RT-qPCR, real-time quantitative PCR; RXR, retinoid X receptor; S5P, serine 5 phosphorylated; shRNA, small hairpin RNA; TF, transcription factor; TH, thyroid hormone; TR, TH receptor; TRPM3, transient receptor potential cation channel M3; TSS, transcription start site.
TH and GC act via ligand-activated transcription factors (TFs) (nuclear hormone receptors [NRs]) (11). Synergistic interactions among TFs during animal development help to establish cellular competency, or may enhance sensitivity to developmental signals that vary in space and time (12).

We recently found that the Krüppel-like factor 9 (Klf9) gene is synergistically transactivated by TH plus GC (see also Ref. 13), it is a direct genomic target for TH receptor (TR) and GC receptor (GR), and it exhibits strong developmental and hormone-dependent regulation in Xenopus and mouse central nervous systems (14–25). Klf9 mediates TH actions on neuronal (15, 16, 26–30) and oligodendrocyte differentiation (28), and Klf9-deficient mice show defects in cerebellar Purkinje cell development, delayed neuronal maturation, impaired neurogenesis-dependent long-term potentiation in the dentate gyrus, and impaired behavior consistent with defects in these brain areas (31, 32). Given its strong hormone regulation, and its demonstrated roles in development, Klf9 may represent a key regulatory node for hormone synergy on phenotypic expression (25).

In the current study we investigated the molecular basis for synergistic transactivation of Klf9 by TH and GC as a model to understand NR synergy. We discovered an ultraconserved enhancer element of approximately 180 bp located 4–6 kb upstream of the transcription start sites (TSSs) of tetrapod Klf9 genes that supports hormone synergy (Klf9 synergy module [KSM]). We show that liganded TR modulates GR recruitment to the KSM and that combined hormone treatment increases serine 5 phosphorylated (S5P) RNA polymerase II (pol II) at the KSM, which leads to the production of enhancer RNAs (eRNAs). We also found that the KSM interacts with the Klf9 promoter and far upstream regions through chromosome looping, and these interactions may be important for KSM enhancer function.

Materials and Methods

Animal care

Early prometamorphic Xenopus tropicalis tadpoles at Nieuwkoop-Faber (33) stages 52–54 were obtained from in-house spawns and maintained in 150-L holding tanks at 23°C on a 12-hour light, 12-hour dark photoperiod. For experiments, tadpoles were transferred to 4-L aquaria (6 tadpoles/aquarium at a density of 2 tadpoles/L; Aquatic Habitats) and treated with vehicle (0.005% dimethylsulfoxide [DMSO] + 0.001% ethanol [EtOH]) or hormones added directly to the aquarium water: T3 (1nM, 5nM, or 50nM; T2877; Sigma) and corticosterone (CORT) (100nM; C2505; Sigma). Hormones were replenished at 12 hours and treatment continued for an additional 12 hours. Animals were euthanized by immersion in 0.1% benzocaine (Sigma), the brains microdissected, and the region of the diencephalon was collected for RNA isolation and gene expression analysis by real-time quantitative PCR (RT-qPCR) (see below).

We purchased adult wild-type C57/B6J mice from The Jackson Laboratory, maintained them on a 12-hour light, 12-hour dark photoperiod, provided food and water ad libitum, and bred them in the laboratory to generate offspring for analysis. We administered hormone injections and then conducted gene expression analysis by RT-qPCR, or chromatin immunoprecipitation (ChIP) assays (described below) on microdissected hippocampal region of the brain. For gene expression analysis, we administered intraperitoneal (ip) injections at postnatal day (PND) 6 of vehicle (corn oil + EtOH), T3 (25 μg/kg BW) (17), CORT (14 mg/kg BW) (18), or T3 plus CORT. The T3 and CORT were first dissolved in EtOH at high concentration and then added dropwise to corn oil with vigorous mixing (final concentration 0.05% EtOH). A separate group of control animals was not injected and not handled. One hour after injection, animals were killed and brains collected for gene expression analysis. For TR ChIP assay we gave ip injections to PND5 mice of vehicle (0.9% saline, 0.05% EtOH) or T3 (25 μg/kg BW) and collected brains 4 hours after injection (17). For GR ChIP assay 1 group of PND6 mice was unhandled (unstressed control) and another received ip injections of CORT (14 mg/kg BW) and killed 4 hours later (18). All animal experiments were conducted in accordance with the guidelines of the University Committee on the Care and use of Animals at the University of Michigan.

RNA extraction, reverse transcription and RT-qPCR

We extracted total RNA from tadpole and mouse brain, and tissue culture cells using TRIzol reagent (Invitrogen). One microgram of total RNA was used for cDNA synthesis using the High Capacity Reverse Transcription kit with ribonuclease inhibitor from Applied Biosystems (Life Technologies Corp). For quantitative PCR we designed TaqMan assays and conducted RT-qPCR using an ABI 7500 fast real-time PCR machine with Absolute qPCR low ROX Mix (ABgene). The primers and TaqMan probes used to analyze Klf9 and Med1 mRNAs spanned exon/intron boundaries (Supplemental Table 1). Standard curves were constructed using cDNA pools to compare relative expression levels. Klf9 mRNA levels were normalized to Gapdh mRNA in mouse cells (Applied Biosystems assay 4331182) or to ribosomal protein L8 (rpl8) in frog cells (34). The Gapdh or rpl8 mRNA levels did not change with any of the treatments (data not shown). To measure Klf9 heterogeneous RNA (hnRNA) or enhancer RNA we treated total RNA with DNase I (20 U; Roche) before cDNA synthesis and designed TaqMan assays to target the single Klf9 intron or the KSM, respectively (Supplemental Table 1). In all cases, we confirmed that RT- qPCR without reverse transcriptase did not yield amplicons (data not shown). To determine the direction of Klf9 eRNA synthesis, strand-specific primers (Supplemental Table 1) were used for cDNA synthesis followed by semiquantitative PCR (Supplemental Materials and Methods).

Plasmid constructs

One-kilobase fragments from 0 to ~7 kb upstream of the X. tropicalis Klf9 TSS were PCR amplified from genomic DNA (Supplemental Table 2). We also PCR amplified the evolutionarily conserved upstream regions of the mouse and frog Klf9.
gene (KSM, described in Results; mouse: 179 bp from −5333 to −5154 bp; frog: 164 bp from −6000 to −5836 bp upstream of TSS). The PCR fragments were cloned into the pGLa.23 promoter-luciferase reporter vector at the XhoI and HindIII sites to generate plasmids containing the mouse or frog KSM region. Single or multiple nucleotide substitutions within the predicted NR half sites (HSs) (see below) of the mouse KSM were introduced using the QuikChange Site-Directed Mutagenesis kit (Stratagene) following the manufacturer’s protocol.

Cell culture and transient transfection assays

Xenopus laevis embryonic fibroblast (XTC-2) and rat epitheli-um (A6)-derived cell lines were cultured at 25°C under an atmosphere of 5% CO2 in Leibovitz L-15 medium (diluted 1:1.5) supplemented with sodium bicarbonate (2.2 g/L), penicillin G (100 U/mL), streptomycin sulfate (100 μg/mL), and 10% fetal bovine serum (FBS). Before initiating hormone treatments, the growth medium was changed to L-15 medium with FBS that had been stripped of thyroid (35) and steroid hormones (36).

The HT-22 cell line was derived from mouse hippocampus and immortalized with the Simian virus 40 T antigen (37, 38). This line exhibits properties of differentiated neurons, eg, they express neuron specific markers such as enolase and the neurofilament triplet but not the glial cell marker, glial fibrillary acidic protein (37–40). It expresses mRNAs for GR and MR, but only the GR is translated into a functional protein (18). We cultured HT-22 cells at 37°C under an atmosphere of 5% CO2 in DMEM (Invitrogen) containing sodium bicarbonate (2.2 g/L), penicillin G (100 U/mL), streptomycin sulfate (100 μg/mL), and 10% fetal bovine serum (FBS). Before initiating hormone treatments, the growth medium was changed to DMEM with hormone-stripped FBS.

For gene expression analysis we seeded cells at a density of 2.5 × 10⁴ cells per well in 6-well plates. When cells reached 80% confluence, and immediately before hormone treatments, we replaced the growth medium with serum-free L-15 (XTC-2 and A6) or DMEM (HT-22). The T3 was dissolved in DMSO and CORT in 100% EtOH and added to the medium to the final concentrations indicated below. Control treatments received an equivalent concentration of vehicle (0.03% DMSO, 0.001% EtOH). To induce nuclear factor κB (NFκB) signaling, we treated HT-22 cells with 1 mg/mL of lipopolysaccharide (LPS) (L3012; Sigma) for 20 hours and then treated with T3 (30nM), CORT (100nM), or T3 plus CORT. All hormone treatments were continued for 4 hours before cell harvest for RNA extraction or dual luciferase assay. Gene expression experiments were repeated at least 3 times with 3–6 replicate wells per treatment.

For luciferase reporter experiments we seeded cells at 6.5 × 10⁴ cells per well in 24-well plates in growth medium and trans- fected at 50%–60% confluence with 250 ng of the pGL4.23 reporter constructs plus 20 ng of pRenilla plasmid using the FuGENE 6 transfection reagent (Roche) following the manufacturer’s protocol. Immediately before transfection the growth medium was replaced with medium containing hormone-stripped FBS without penicillin/streptomycin, and cells were incubated with the transfection mixture overnight. Cells were then treated in serum-free medium with vehicle (0.03% DMSO, 0.001% EtOH), T3 (30nM), CORT (100nM), or T3 plus CORT for 4 hours before harvest for luciferase activity using the Dual Luciferase Assay System (Promega). Firefly luciferase activity was quantified using a luminometer (Femtometer FB 12; Zylux Corp) and normalized to Renilla luciferase activity. Each transfection experiment was conducted at least 3 times with 4–5 wells per treatment.

Electrophoretic mobility shift assay

We conducted EMSA as described by Hoopfer et al (24) with recombinant X. laevis TRβ and retinoid X receptor (RXRα), and human GR that we synthesized in vitro using the TnT SP6 quick-coupled translation system (Promega). We programmed the reactions with 1 μg of pSP76-xTRB, pSP76-xRXRα (gifts of Yun-Bo Shi) or CMV-hGR (Origene) plasmids and confirmed protein expression by [35S]-labeled amino acid incorporation, SDS-PAGE, and autoradiography (24). Wild-type and mutant duplex oligonucleotides corresponding to the predicted direct repeat with 4-base spacer (DR+4) T3 response element (T3RE), the GRE/MRE (at −5.3 kb from the TSS) (18), and predicted HSs 3–5 located within the mouse KSM, and the upstream GRE/ MRE (at −6.1 kb from the TSS) (18) (Supplemental Table 2) were labeled with 32P-dCTP by Klenow fill-in for use as probes in EMSA.

ChIP assay

We isolated chromatin from HT-22 cells treated with vehicle (EtOH + DMSO), T3 (30nM), CORT (100nM), or T3 plus CORT. We also isolated chromatin from the hippocampal region of PND6 mice 1 hour after injection of vehicle (oil), or CORT (14 mg/kg); a group of control animals was left unhandled. Five micrograms of sheared chromatin were used for each reaction, and ChIP assays were conducted as described by Denver and Williamson (17). We used a rabbit polyclonal anti-serum raised against the full-length X. laevis TRβ (PB antisem- rum, 5 μL; it does not distinguish TRα from TRβ; provided by Yun-Bo Shi). This antiserum has been used extensively for ChIP assay on frog tissues (41–43) and in mouse (17, 44). The frog and mouse TR proteins share greater than 90% sequence identity. We also used commercial antibodies to mouse GR (5 μg; M-20X; Santa Cruz Biotechnology, Inc), RNA pol II C-terminal domain (2 μg; 5408; Abcam), S5P pol II (2 μg; 5131–50; Abcam), S2P pol II (2 μg; 5095; Abcam), NFκB p50 subunit (p50) (3 μg; 7971; Abcam), histone 3 (H3) (5 μL; 06–753; Millipore), acetylated H3 (acH3) (5 μL; 06–599 Millipore), acH4 (5 μL; 06–866; Millipore), or normal rabbit serum (NRS) (Sigma). We also used ChIP samples that were prepared previously by Den- ver and Williamson (17) from PND5 vehicle or T3-injected mouse brain, and vehicle or T3-treated N2a[TRβ] cells. We analyzed ChIP DNA by quantitative real-time PCR using Taq-Man primer/probe sets that targeted different regions of the mouse Klf9 gene (Supplemental Table 1). Each of the TaqMan primer/probe sets had similar PCR efficiency.

Chromosome interaction analysis by paired end tag sequencing (ChIA-PET)

The ChIA-PET method allows for determination of protein- DNA binding sites and the physical interactions between nu- clear proteins supported by chromosomal looping (45, 46). We treated Nieuwkoop-Faber stage 50 X. tropicalis tadpoles with 10nM T3 for 24 hours, isolated chromatin from a pool of 8 tail fins, and conducted ChIP and ChIA-PET following our pub- lished methods (45, 47–49). Briefly, long-range chromatin in- teractions were captured by DNA-protein cross-linking with formaldehyde. Chromatin was fragmented by sonication, and
DNA-protein complexes were enriched by ChIP using polyclonal antiserum against X. tropicalis TR. Tethered DNA fragments were connected with DNA linkers by proximity ligation. The ligation generated 2 types of products: self-ligation of the same DNA fragments and interligation between different DNA fragments in individual chromatin complexes. Short tag signature information (20–30 bp) from the 2 ends of target DNA fragments (PETs) was extracted by restriction enzyme (RE) digestion. The PETs were sequenced on the SOLiD platform. The resulting ChIA-PET sequences were mapped to the reference X. tropicalis genome. Self-ligation PET sequences were mapped in the reference genome within a 3-kb span, demarcating ChIP DNA fragments, similar to the standard ChIP-sequencing method. Interligation PET sequences reveal close spatial proximity through chromatin looping. Singleton PETs are presumed to be experimental noise, and overlapping PET clusters are considered enriched putative binding sites or interaction events. Full details of the experimental procedures and the complete datasets generated by ChIA-PET applied to X. tropicalis will be published elsewhere.

**Chromosomal conformation capture (3C)-qPCR assay**

We conducted 3C-qPCR assay on chromatin isolated from HT-22 cells as described by Hagège et al (50) with minor modifications. We plated HT-22 cells in 10-cm dishes in DMEM with 10% hormone-stripped FBS. After 48 hours, cells were treated with 30nM T3 plus 100nM CORT for 4 hours, and single cell suspensions were prepared as previously described (50). Cross-linking of protein and DNA was done by gently tumbling cells in 10-mL PBS with 10% hormone-stripped FBS and 2% formaldehyde at room temperature for 10 minutes. A second protein-protein cross-linking step was conducted by tumbling cells in 10-mL PBS with 10% hormone-stripped FBS and 0.2mM dithiothreitol. Genomic DNA was isolated, digested with restriction enzymes HinII or HinI1003, ligated, and purified by phenol-chloroform extraction and EtOH precipitation.

As a control for template and primer efficiency, a bacterial artificial chromosome clone containing the mouse Klf9 locus was similarly digested with DpnII, ligated and purified by phenol-chloroform extraction. Sample purity and loading adjustments for qPCR were conducted as described in Hagège et al (50).

Positive interactions, as indicated by the presence of ligated chromatin fragments, were detected by the presence of qPCR amplicons (80–120 bp in length) using a constant reverse primer and a TaqMan probe targeted to the mouse Klf9 promoter, and forward primers targeted within 50 bp of DpnII cutting sites at several genomic region covering 10 kb upstream of the TSS (Supplemental Table 3). The HindIII-digested and ligated chromatin fragments served as a negative control for qPCR amplification because the combination of different forward primers (designed close to a DpnII cutting site) and the constant reverse primer and TaqMan probe, should yield amplicons more than 800 bp, which are beyond the limit of detection of the qPCR assay. We express the data based on positive interactions for 8 independent samples (4 vehicle and 4 hormone-treated samples) as detected by the presence of amplification products by qPCR.

**Data analysis and statistics**

Data were analyzed by one-way ANOVA or Student’s t test using SYSTAT (α < 0.05; version 10, SPSS Inc). Derived values from dual luciferase and ChIP assays were log_{10} transformed before statistical analysis. A synergistic effect of T3 and CORT on gene expression and ChIP signal is defined as: 1) a greater than additive effect by one standard deviation with combined hormone treatment, or 2) no effect with T3 or CORT alone but a significant effect with combined hormone treatment (2). To predict putative TF binding sites we used the online programs Transcription Element Search System (www.cbil.upenn.edu/tess) and Match (www.gene-regulation.com/pub/programs.html#match) using a library of matrices from TRANSFAC 6.0 (www.gene-regulation.com) with core and matrix similarity of TF binding sites set higher than 0.85. The conservation of the identified TF binding sites and flanking sequences was assessed using BLAT (http://genome.ucsc.edu/). Extracted sequences were aligned using Clustal W. The percent similarity to human was calculated for the T3,RE and GRE/MRE HS, and nucleotide frequency diagrams were produced with Weblogo (http://weblogo.berkeley.edu/logo.cgi).

**Results**

**Synergistic transactivation of the Klf9 gene by T3 plus CORT**

Treatment of XTC-2 or A6 cells with T3 or CORT for 4 hours increased Klf9 mRNA above control, whereas combined hormone treatment caused synergistic increases, with the largest effect (53-fold increase) seen in A6 cells (Figure 1A). In HT-22 cells T3 and CORT also independently and synergistically increased Klf9 mRNA.
The synergistic effect was seen at 1 hour and persisted through 6 hours of hormone treatment (Figure 1B).

We also saw hormone synergy on Klf9 hnRNA in HT-22 cells, supporting increased gene transcription (Supplemental Figure 1). Treatment with the protein synthesis inhibitor cycloheximide (CHX) (Supplemental Materials and Methods) did not affect hormone-dependent induction of Klf9 mRNA in either XTC-2 or HT-22 cells which also supports direct hormone action on Klf9 gene transcription (Supplemental Figure 2).

We next investigated whether hormone synergy occurs on Klf9 in tadpole and mouse brain in vivo. Treatment with T3 or CORT alone increased Klf9 mRNA in tadpole brain, whereas combined hormone treatment caused a synergistic increase (Figure 1C). In mouse, ip injection of T3 or CORT alone increased Klf9 mRNA in the hippocampal region of the brain 1 hour after injection, whereas combined hormone treatment caused a synergistic increase (Figure 1D).

**An evolutionarily conserved approximately 180-bp genomic region located upstream of tetrapod Klf9 genes supports synergistic transactivation by T3 plus CORT**

To identify genomic regions that confer hormone-dependent synergistic transactivation of Klf9, we used promoter-reporter assays in XTC-2 cells to screen 1-kb fragments of the X. tropicalis Klf9 5′-flanking region covering 0 to −7 kb upstream of the TSS. We found that the fragment located −5 to −6 kb upstream of the TSS was T3 and CORT-responsive, and this region supported synergistic transactivation (Figure 2A; see also Supplemental Figure 3 for HT-22 cells).

Bioinformatics analysis revealed an approximately 180-bp segment within the −5- to −6-kb fragment that is highly conserved among tetrapod Klf9 genes (Figure 2B and Supplemental Tables 4 and 5). We found orthologous sequences in most tetrapod genomes, and comparative sequence analysis showed that this intergenic region is remarkably conserved among tetrapods, eg, sequence similarity was 100% between human and chimp and 75% between human and frog (Supplemental Table 6). Sequence similarity within the coding regions of tetrapod Klf9 genes was sim-
Figure 2. Identification of an evolutionarily conserved approximately 180-bp genomic region upstream of tetrapod Klf9 genes that supports synergistic transactivation by TH and CORT. A, Enhancer-reporter scan of the 5′-flanking region of the X. tropicalis Klf9 gene; 1-kb fragments covering 0 to −7 kb upstream of the Klf9 TSS were cloned into the luciferase vector pGL4.23 and transfected into XTC-2 cells. We treated cells with vehicle (0.03% DMSO, 0.001% EtOH), T3 (5nM), CORT (100nM), or T3 plus CORT for 4 hours before harvest and analysis by dual luciferase assay. The −5- to −6-kb fragment supported individual hormone responses and hormone synergy (F(3,15) = 89.44, P < .001; ANOVA). The −6- to −7-kb fragment showed a small response to CORT (F(3,15) = 11.64, P = .05; ANOVA) that was unaffected by T3. Bars represent the mean ± SEM (n = 4–5/treatment) fold change, and letters above the means indicate significant differences among hormone treatments within a promoter fragment (means with the same letter are not significantly different; Tukey’s multiple comparison test; P < .05). The dashed line indicates the calculated additive effect of combined hormone treatment. B, A genomic region spanning approximately 180 bp upstream of tetrapod Klf9 genes is evolutionarily conserved (the KSM). Shown is the most highly conserved region spanning 138–146 nt for 5 tetrapods (see Supplemental Tables 4 and 5 for more comprehensive analyses). The highlighted sequences show the highly conserved GRE/MRE and DR4T3RE regions. The evolutionarily conserved, predicted NFkB site that overlaps with the T3RE is indicated by a bar over the sequence. Note that there is a 2-base insertion in nonmammalian tetrapods. Genomic location of the KSM relative to the TSS for species for which it was possible to determine: X. tropicalis: −6000 to −5836 bp; mouse: −5333 to −5154 bp; human: −4634 to −4505.

Figure 2. Identification of an evolutionarily conserved approximately 180-bp genomic region upstream of tetrapod Klf9 genes that supports synergistic transactivation by TH and CORT. A, Enhancer-reporter scan of the 5′-flanking region of the X. tropicalis Klf9 gene; 1-kb fragments covering 0 to −7 kb upstream of the Klf9 TSS were cloned into the luciferase vector pGL4.23 and transfected into XTC-2 cells. We treated cells with vehicle (0.03% DMSO, 0.001% EtOH), T3 (5nM), CORT (100nM), or T3 plus CORT for 4 hours before harvest and analysis by dual luciferase assay. The −5- to −6-kb fragment supported individual hormone responses and hormone synergy (F(3,15) = 89.44, P < .001; ANOVA). The −6- to −7-kb fragment showed a small response to CORT (F(3,15) = 11.64, P < .05; ANOVA) that was unaffected by T3. Bars represent the mean ± SEM (n = 4–5/treatment) fold change, and letters above the means indicate significant differences among hormone treatments within a promoter fragment (means with the same letter are not significantly different; Tukey’s multiple comparison test; P < .05). The dashed line indicates the calculated additive effect of combined hormone treatment. B, A genomic region spanning approximately 180 bp upstream of tetrapod Klf9 genes is evolutionarily conserved (the KSM). Shown is the most highly conserved region spanning 138–146 nt for 5 tetrapods (see Supplemental Tables 4 and 5 for more comprehensive analyses). The highlighted sequences show the highly conserved GRE/MRE and DR4T3RE regions. The evolutionarily conserved, predicted NFkB site that overlaps with the T3RE is indicated by a bar over the sequence. Note that there is a 2-base insertion in nonmammalian tetrapods. Genomic location of the KSM relative to the TSS for species for which it was possible to determine: X. tropicalis: −6000 to −5836 bp; mouse: −5333 to −5154 bp; human: −4634 to −4505.

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The Endocrine Society. Downloaded from press.endocrine.org by [individualUser.displayName] on 09 June 2015 at 11:34. For personal use only. No other uses without permission. All rights reserved.
Mutation of both HSs of the predicted T₃RE (HS4+HS5 mut) eliminated all hormone responses.

**TR and GR bind to the mouse KSM in vitro**

Using EMSA we found that TR-RXR heterodimers bound a probe containing the predicted T₃RE within the mouse KSM (Figure 4A). Mutation of the 5’ T₃RE HS (HS4) reduced, whereas mutation of both HSs eliminated TR-RXR binding (Figure 4A). Because the KSM reporter with a mutated 3’ T₃RE HS (HS5) retained some T₃ responsiveness (and hormone synergy; data not shown), we investigated whether TR-RXR could bind a probe containing only the predicted HSs HS3 and HS4. However, TR-RXR was unable to bind this DNA sequence (Figure 4A).

The GR bound the previously characterized −5.3-kb GRE/MRE (HS1 and HS2) within the KSM, which was competed for by wild-type but not mutant HS1+HS2 radioinert oligonucleotide (Figure 4B) (18). Because the KSM reporter with mutations in both GRE/MRE HSs retained some CORT responsiveness (see above) we investigated whether GR could bind a probe containing HS3–HS5. However, the GR was unable to bind this DNA sequence (18).

In silico analysis identified other TF binding sites in the mouse KSM and at the region of the upstream GRE/MRE (−6.1 kb) and T₃RE (−6.3 kb), which included consensus NFκB sites (18), one of which overlaps with the T₃RE within the KSM (conserved among mammals but with a 2-base insertion in other tetrapods) (Figure 2B and Supplemen tal Figure 7; see also Supplemental Tables 7 and 8). Using ChIP assay, we found that p50 associated with these regions in HT-22 cell chromatin, and the signal was decreased by treatment with CORT (Supplemental Figure 8A); we did not detect p50 at the mouse Klf9 intron. Activation of NFκB signaling by LPS enhanced the Klf9 mRNA response to T₃ in HT-22 cells (Supplemental Figure 8B).

**TR and GR associate with the KSM in mouse brain chromatin**

ChIP assay for TR on chromatin isolated from the hippocampal region of PND6 mice 1 hour after ip CORT injection showed CORT-dependent association of GR at the KSM and also at the previously identified GRE/MRE-6.1 located upstream of the KSM (found only in therian mammals) (Figure 4D) (18). By comparison, we detected no significant GR ChIP signal at the distal intron of the mouse Klf9 gene.

**Liganded TR enhances CORT-dependent recruitment of GR, but not TR, across the mouse Klf9 locus**

Analysis of TR and GR association across the Klf9 locus in HT-22 cells by ChIP assay showed a strong TR signal only at the KSM, which was unaffected by hormone treatment, consistent with TRs being constitutively bound to DNA (Figure 5). We did not see TR association in HT-22 cells at the −3.8-kb T₃RE region that we previously identified in mouse brain and Neuro2a[TRβ1] cells (Figure 4C; although we saw strong TR signal at the KSM in Neuro2a[TRβ1] cells (17 and data not shown). Also, we did not see TR signal in HT-22 cells at the −6.2-kb T₃RE identified by Chatonnet et al (19) in the mouse cerebellar-derived cell line C17.2 engineered to overexpress TRα1 or TRβ1. Thus, mammalian Klf9 genes have T₃REs located outside the KSM, which may be used in a tissue and/or developmental-stage dependent manner as compared with the ubiquitous function of the T₃RE within the KSM. The ChIP signal for the NRS control was low to nondetectable across the Klf9 locus and was not affected by hormone treatment (data not shown).

The GR ChIP signal was low across the Klf9 locus in HT-22 cells and was not affected by T₃ (Figure 5B). There was a trend towards higher GR signal at the KSM after exposure to CORT but this did not reach statistical significance (P = .073). However, combined treatment with T₃ plus CORT caused a robust increase in GR ChIP signal at the KSM and at the −6.1-kb GRE/MRE (Figure 5).

Our GR ChIP assays support that liganded TR promotes recruitment of liganded GR to the Klf9 locus, thereby enhancing CORT-dependent transcription of the Klf9 gene. This may represent a key molecular event in the synergistic transcriptional response to hormones. Consistent with this hypothesis, we found that forced expression of a dominant negative TR (dnTR) that does not bind hormone in both HT-22 and XTC-2 cells reduced CORT-dependent induction of Klf9 mRNA (also reduced the T₃ response and hormone synergy) (Supplemental Materials and Methods and Supplemental Figure 9).
Figure 4. TR and GR bind in vitro to predicted hormone response elements and associate in chromatin with Klf9 upstream regions in mouse hippocampus. A, EMSA for TR-RXR binding to [32P]-labeled oligonucleotide probes (Supplemental Table 2) corresponding to wild-type (wt) or mutated (mt) hormone response elements located upstream of the mouse Klf9 gene (shown above the graphs). Probes were incubated with or without in vitro synthesized TR/H9252 plus RXR/H9251 before electrophoresis. The arrow indicates supershifted bands due to TR-RXR binding to the radiolabeled probes. The TR-RXR showed strong binding to the wtHS4 wtHS5 probe (HS, within the KSM shown in the diagram above the graphs), which corresponds to the predicted DR4T3RE located within the KSM, but not to other probes. B, Competitive EMSA for GR binding to [32P]-labeled oligonucleotide probes (Supplemental Table 2) corresponding to wt hormone response elements located at the mouse KSM (at ~5.3 kb upstream of the TSS) and the GRE/MRE located upstream of the KSM (at ~6.1 kb upstream of the TSS; identified by Bagamasbad et al [18]). Probes were incubated with or without in vitro synthesized GR before electrophoresis. In lanes 3 and 4, the reactions were incubated with 1.5 µM radioinert competitor: 1) wtHS1 + wtHS2; 2) mtHS1 + mtHS2. The arrows indicate supershifted bands due to GR binding to the radiolabeled probes. The GR bound to the GRE/MRE-5.3 kb and GRE/MRE-6.1 kb probes but not to the wtHS3–5 probe; GR binding to the GRE/MRE-5.3 kb probe was competed by wt but not by mutant radiolabeled oligonucleotide. C, TR association at the Klf9 locus in mouse hippocampus analyzed by ChIP assay. We administered ip injections of vehicle or T3 (25 µg/kg BW; n = 4–5/treatment) to wt PND5 mice and dissected the hippocampal region 4 hours later for chromatin extraction for TR ChIP assay. We analyzed ChIP DNA by RT-qPCR using TaqMan assays that targeted the region of the predicted T3RE at ~6.2 kb from the Klf9 TSS, the T3RE within the KSM (~5.23 kb) and a distal intronic region (~11 kb; negative control region) of the mouse Klf9 gene. Bars represent the mean ChIP signal expressed as a percentage of input for TR serum (anti-TR) or NRS (negative control). The asterisks indicate statistically significant differences between the anti-TR and NRS groups (*, P < .05; **, P < .01; ***, P < .001; Student’s 2 sample t test; derived values were log10 transformed before statistical analysis). D, GR association at the Klf9 locus in mouse hippocampus analyzed by ChIP assay. wtPND6 mice were left unhandled or given ip injections of CORT (14 mg/kg BW; n = 9/treatment). One hour after injection, animals were killed, the hippocampal region was dissected, and chromatin was extracted for GR ChIP assay. The ChIP DNA was analyzed by RT-qPCR using TaqMan assays that targeted the GRE/MRE ~6.1 kb, the GRE/MRE at ~5.3 kb within the KSM, and a distal intronic region (~11 kb; negative control region) of the mouse Klf9 gene. Bars represent the mean ChIP signal expressed as a percentage of input. Asterisks indicate statistically significant differences between unhandled and CORT-injected animals (P < .05; Student’s 2 sample t test; derived values were log10 transformed before statistical analysis).
Recruitment of RNA pol II across the Klf9 locus

ChIP assays conducted on chromatin from HT-22 cells showed association of pol II across the mouse Klf9 locus, but the total pol II signal did not vary by genomic region. We also looked at the association of 2 phosphorylated forms of pol II, S5P, which is involved with transcription initiation, and S2P, which functions in transcription elongation. The mouse KSM is bidirectionally transcribed to generate long noncoding RNAs (lncRNAs).

The presence of pol II at the KSM prompted us to investigate whether this genomic region is transcribed into RNA. We detected transcripts at the KSM in HT-22 cells and mouse hippocampus (Figure 6; we refer to this RNA as the Klf9 eRNA). The estimated size of this eRNA is between 1.3 and 4 kb, and it is bidirectionally transcribed (Supplemental Figure 11). The UCSC Genome Browser shows a lncRNA associated with the mouse KSM predicted by Ensembl (54).

In HT-22 cells, the pattern of hormone response of the Klf9 eRNA was similar to that of the mRNA, but the magnitude of increase in Klf9 eRNA after combined hormone treatment was significantly greater than the Klf9 mRNA (Figure 6A; see also Supplemental Figure 11). Similar to the mRNA, the level of Klf9 eRNA was not affected by protein synthesis inhibition with CHX (Figure 6B). In hippocampus of PND6 mice we saw significant increases in Klf9 eRNA after T3 or T3 plus CORT injection (not synergy) but not with CORT alone (F(4,17) =

Hormone treatment causes nucleosome repositioning and histone hyperacetylation at the KSM in HT-22 cells

ChIP assays for H3, acH3, and acH4 showed that T3 or CORT decreased levels of H3 but increased acH3 (normalized to H3) and acH4 (normalized to H3) at the KSM (Supplemental Figure 10) and −6.1-kb GRE/MRE (data not shown). There was no further enhancement by combined hormone treatment at either genomic region. Hormone treatment did not alter H3, acH3, or acH4 levels at the distal intron (+11 kb from the TSS; data not shown).

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Figure 5. Hormone-dependent recruitment of TR, GR, and S5P (initiating) and S2P (elongating) RNA pol II across the mouse Klf9 locus. We treated HT-22 cells with vehicle, T3 (30nM), CORT (100nM), or T3 plus CORT for 4 hours before harvest for chromatin extraction and ChIP assays. We analyzed ChIP samples (n = 6/treatment) by RT-qPCR using TaqMan assays that targeted different genomic regions of the mouse Klf9 gene (shown on the x axis, with distances relative to the Klf9 TSS). Shaded bars highlight the KSM and the TSS. A, Strong TR ChIP signal was seen only at the KSM, but unaffected by hormone treatment. B, The GR ChIP signal was not affected by T3; CORT increased the mean GR signal, but this did not reach statistical significance (P = .073). Combined hormone treatment increased GR ChIP signal at the KSM (F(3,21) = 8.631, P < .001; ANOVA) and at the GRE/MRE-6.1 kb (F(3,21) = 7.4555, P < .05). C, The S5P pol II ChIP signal peaked at the KSM, −3.8 kb from the TSS, and at the TSS. There were no effects of individual hormone treatments but combined hormone treatment increases S5P pol II signal at the KSM and −3.8 kb (KSM: F(3,18) = 10.6, P < .001; −3.8 kb: F(3,20) = 2.43, P < .05; ANOVA). T3 or T3 plus CORT increased the S5P pol II signal at the TSS, but CORT alone had no effect (F(3,18) = 11.31, P < .0001). D, The S2P pol II ChIP signal peaked at the KSM was increased by all hormone treatments, and was highest after combined hormone treatment (F(3,21) = 6.736, P < .01). All hormone treatments similarly increased S2P pol II signal at −3.8 kb (F(3,18) = 3.8, P = .028). Each point represents the mean, and means with the same letter are not significantly different (ANOVA, Tukey’s multiple comparison test; P < .05). Absence of letters indicates that there were no statistically significant differences among treatments at that genomic location. Statistical analysis was conducted on log10-transformed data.

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of the Klf9 gene. We demonstrated interactions between the mouse KSM core promoter and genomic regions that span the KSM (~4.9 kb from the TSS) (Figure 7B), which were not influenced by hormone treatment (data not shown). We detected no amplicons when the same primer-probe pairs were used with HindIII-digested and religated samples (negative ligation controls; data not shown).

Both the frog and human ChIA-PET datasets also identified far upstream regions with which the KSM interacts (Figure 7A and Supplemental Figure 13). In the frog this region is located at ~39.2 kb from the TSS, in human at ~72 kb and in mouse at ~90 kb. There is clear conservation across several primates that had sequences available for this far upstream interacting region, and there are homologous regions in some other eutherian mammals (Supplemental Table 6). The frog and mammalian upstream interacting regions do not appear to be conserved. In human the ~72-kb region has a strong peak for H3 lysine 4 (K4) monomethylation (Supplemental Figure 13), a mark of active enhancers (56).

Knockdown of Med1 in HT-22 cells reduced hormone induction of Klf9 mRNA and eRNAs

The Mediator complex subunit MED1 is important for linking enhancers to gene promoters by chromosomal looping (57–59) and associates with TR (60) and GR (61, 62) in a ligand-dependent manner. Subunits of the Mediator complex MED1 and MED12 exhibit distinct peaks that are associated with peaks of pol II at the mouse KSM and promoter, and at 4 upstream regions (Supplemental Figure 14 and Supplemental Table 6) (57). Transduction of HT-22 cells with Med1 (Supplemental Materials and Methods) caused an approximately 60% reduction in Med1 mRNA levels compared with the scrambled shRNA control (Supplemental Figure 15). Knockdown of Med1 did not alter baseline Klf9 mRNA but caused small but statistically significant reductions in the responses to T3 and T3 plus CORT. Baseline Klf9 eRNA levels were similarly unaffected by Med1 shRNA, but hormone re-
responses were significantly reduced (Supplemental Figure 15). The relatively small reductions in the T3-induced Klf9 mRNA levels may be due to the incomplete knockdown of Med1.

Discussion

Cooperativity among TFs provides a means for cells to integrate multiple inputs to regulate gene transcription (12, 63, 64). Synergistic transcriptional responses, in part due to TF cooperativity, allow for 2 or more simultaneous signals to generate a response greater than additive, or for one signal to increase the sensitivity of the cell to a second, perhaps temporally delayed signal. Here, we report synergistic regulation of the Klf9 gene by T3 and CORT that may be explained by the presence of an evolutionarily conserved NR enhancer module located in the 5′-flanking region of the gene. This element (the KSM) contains a T3RE that overlaps with a NFκB site and a GRE/MRE in close proximity. We show that liganded TR modulates recruitment of liganded GR and SP5 pol II to the locus, and together, liganded TR and GR generate a synergistic transcriptional response. The region of the KSM is transcribed to generate eRNAs which are also synergistically induced by hormone action, and the production of eRNAs may depend on the loading of pol II from the promoter onto the KSM, or vice versa, requiring chromosomal looping between these genomic regions, mediated in part by the Mediator complex (57, 59, 65, 66).

An ultraconserved genomic region supports hormone synergy on Klf9 gene transcription

Earlier findings from our group and others showed that Xenopus and mouse Klf9 genes are regulated by T3.
and CORT (15–18). Gil-Ibánez et al (13) recently reported that treatment of primary cultures of cerebrocortical cells from wild-type mice with T3 plus dexamethasone caused a synergistic increase in Klf9 mRNA (see also Ref. 67). Here, we show that frog and mouse Klf9 genes are synergistically induced by combined hormone treatment, further supporting that hormone regulation of this gene arose early in the evolution of tetrapods and has been conserved. An enhancer-reporter scan of the 5’-flanking region of the frog Klf9 gene, and comparative genome sequence analysis revealed an ultraconserved (68, 69) approximately 180-bp sequence located 4–6 kb upstream of tetrapod Klf9 genes that supports hormone synergy. It is located between the Klf9 and TRPM3 genes in tetrapod genomes. We could not find orthologous regions in other chordates, supporting that the KSM evolved in tetrapods. The KSM exhibits higher sequence conservation than the core promoter of Klf9 genes, suggesting that it has been “ultraselected” (70).

Located within the KSM are 5 predicted hexanucleotide HSs (HS1–HS5) that are completely conserved across tetrapods and could therefore mediate NR actions. Our previous work showed that the GRE/MRE at −5.3 kb (HS1 and HS2) in the mouse KSM is a bona fide GRE/MRE (18). The HS4 and HS5 were previously identified by Furlow and Kanamori (23) in Xenopus and predicted to function as a DR +4 T3RE in vivo. We confirmed that TR associates in chromatin with the orthologous mouse T3RE at −5.23 kb, and mutational analysis showed that it is required for T3-dependent transactivation. Although mutation of the GRE/MRE in the KSM reduced CORT and T3 responses, mutation of the KSM T3RE eliminated all hormone responses. Together with the observation that expression of a dnTR reduced the CORT response in tissue culture cells, our findings support that the T3RE within the KSM plays a key role in modulating activity of this enhancer, and supports a model whereby liganded TR facilitates entry of GR to the KSM, thus supporting hormone synergy.

Within the KSM there are 2 predicted NFκB sites, one of which overlaps with the 5’ HS of the T3RE. NFκB is comprised of a family of constitutively expressed TFs that are activated by the proinflammatory cytokine TNFα, among other signals, and bind to NFκB response elements of target genes as dimers (p50 and p65 dimers) (71). We found that p50 associates at the KSM and that activation of NFκB signaling by treatment of HT-22 cells with LPS acted synergistically with T3 to induce Klf9 mRNA. Thus, our findings support that TR-RXR and NFκB, possibly acting via a composite response element, cooperate to regulate transcription of Klf9. There are a few reports in the literature of interactions, both positive and negative, between the thyroid axis and inflammatory pathways mediated by NFκB (72–75), but this area requires more investigation.

Analysis of human ENCODE data support that the KSM is in a transcriptionally active chromatin environment that can be modulated by hormones. We found that histone modifications at the mouse KSM are modulated by hormone action. For example, T3 or CORT increased acH3 and acH4, marks of transcriptionally active chromatin (76), and reduced H3 at the KSM, indicative of nucleosome repositioning and chromatin disassembly (77). The responses to combined hormone treatment were similar to the individual hormones, suggesting that although hormone responses are promoted by chromatin decompaction, hormone synergy is not dependent on further enhancement of histone acetylation or nucleosome repositioning (Supplemental Figure 10).

**Enhancer transcription and chromatin interactions**

We found thatpol II is enriched at the KSM, which supports recent findings that the preinitiation complex is recruited to enhancers, and that pol II-bound enhancers may function in promoter targeting and regulation (78). Genome-wide analyses show that a large proportion of pol II, general TF, and chromatin modifying enzyme peaks are found at enhancers (56, 79–83), which is likely to be an important step in gene regulation that maintains the locus in a poised state without the risk of premature gene transcription (78, 84). In our studies, when both TR and GR were liganded, we saw enhanced recruitment of the elongating isoform (S5P) of pol II, which likely supported increased transcription, as evidenced by increased Klf9 hnRNA, mRNA, and also eRNA.

Consistent with the enrichment of pol II at enhancer regions, recent findings show that enhancers are transcribed to give rise to a type of long, intergenic noncoding RNA (eRNA) (81). We detected eRNA transcripts from the region of the mouse and frog KSMs, and several expressed sequence tags associated with the KSM can be found in vertebrate genome databases. The synergistic response to combined hormone treatment was significantly greater for the Klf9 eRNA than for the mRNA. Others have found that eRNAs increase in parallel to the associated protein coding genes in response to stimuli (79, 81). Genome-wide expression analysis of lncRNAs in mouse found that brain lncRNAs are more evolutionary constrained than lncRNAs expressed in other tissues, and tend to be close to genes that code for proteins involved with transcriptional regulation and nervous system development (85). Consistent with these findings, the KSM is highly conserved among tetrapods, Klf9 exhibits strong developmental expression in the central nervous system.
that depends on TH (16, 17, 22, 24), and KLF9 functions in T₃-dependent actions on neuron and oligodendrocyte differentiation (16, 27, 28, 31). We do not know whether the Klf9 eRNA has a biological function, but we speculate that it could participate in communication between the KSM and the Klf9 promoter. Recent findings support that lncRNAs may play a role in regulating gene transcription (78), possibly by influencing chromosomal looping through direct interaction with the Mediator complex, and thereby stabilizing the interaction between enhancer and promoter (59, 85–87).

Using ChIA-PET and 3C-qPCR analysis we found that chromosomal looping brings the KSM into contact with the Klf9 core promoter (and several far upstream regions). Taken together with our functional analyses, these findings support that the KSM is a bona fide enhancer that communicates with the Klf9 promoter to control gene transcription. The Mediator complex subunit MED1 functions in chromosomal looping (57–62) and ChIP sequencing data for MED1 and MED12 conducted on mouse embryonic stem cells (57) showed that both exhibited strong peaks at the KSM, Klf9 promoter, and several upstream regions. Knockdown of MED1 in HT-22 cells reduced T₃-induction of Klf9 mRNA and eRNAs. It is interesting that the eRNA was affected, even more strongly than the mRNA. This suggests that MED1 is important for enhancer transcription, perhaps for loading pol II from the promoter onto the enhancer. This would be consistent with the findings of Kim et al (81), who showed that eRNA synthesis depended on the presence of the promoter of the associated gene.

**Molecular basis for hormone synergy**

There are several proposed molecular mechanisms that could account for transcriptional synergy. For example, there may be enhanced recruitment of chromatin modifying enzymes to the locus (88); recruitment of an accessory TF at regulatory regions that only occurs in the presence of the synergizing NRs/TFs (89, 90); unidirectional or reciprocal cooperative binding of the synergizing NRs/TFs that may be attributed to enhanced decompaction of chromatin, thereby allowing for greater DNA-protein contact (91–96); and interaction between 2 regulatory regions mediated by protein-protein interactions and perhaps chromosomal looping (97, 98). It is likely that several of these mechanisms operate at the Klf9 locus to support hormone synergy.

Our findings support that TRs, which are resident in chromatin in the unliganded state, modulate access of liganded GR to the KSM. We found that T₃ was required for CORT-induced recruitment of GR to the KSM, which also resulted in enhanced S5P pol II signal and strongly increased enhancer transcription. Forced expression of a dnTR unable to bind hormone and recruit coactivators suppressed CORT-mediated Klf9 transactivation, suggesting that GR entry is dependent on chromatin state as modulated by the TR. Our findings also support that chromosomal looping is important for hormone action at the Klf9 locus. Thus, TRs resident in chromatin in the unliganded state may function as “gate keepers” to modulate local chromatin structure (99), and therefore the accessibility of genomic loci to allow for developmental stage and physiological state-appropriate gene expression (100).

**Acknowledgments**

We thank Dr J. David Furlow (University of California, Davis, CA) for providing sequence data for the 2 Xenopus laevis Klf9 genes, to Dr David Schubert (Salk Institute, La Jolla, CA) for providing the HT-22 cells, and Yun-Bo Shi (National Institute of Child Health and Development, National Institutes of Health, Bethesda, MD) for providing plasmids and antiserum. Elysha Cloyd and Daniel Brown provided technical assistance. Dr Kenneth Cadigan, Dr Audrey Seaholtz, and Dr Cunming Duan provided valuable comments on the project while serving on P.B.’s PhD dissertation committee. We also thank Patrice Bilesimo for the production of the TR ChIP material used for ChIA-PET; Zhenshui Zhang, Huay Mei Poh, and Alexis Khng Jiaying for construction of DNA-PET, RNA-PET, and Chia-PET libraries, and the Genome Institute of Singapore GTB sequencing group personnel for Illumina sequencing.

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This work was supported by the National Institute of Neurological Disorders and Stroke, the National Institutes of Health Grant 1 R01 NS046690, and the National Science Foundation Grant IOS 0922583 (to R.J.D.). P.B. was supported by a predoctoral fellowship from the Rackham Graduate School and an Edwin Edwards Scholarship from the Department of Molecular, Cellular and Developmental Biology, University of Michigan. J.R.K. was supported by an MCubed grant from the University of Michigan. L.S. and Y.R. were supported by “CRESCENDO,” an Integrated Project from FP6 (Contract LSHM-CT-2005–018652) and a Merlion Project (Contract 5.01.10). This research used the Cell and Molecular Biology Core of the Michigan Diabetes Research and Training Center funded by NIH P60 DK20572 from the National Institute of Diabetes and Digestive Kidney Diseases and the Vector Core at the University of Michigan.

Disclosure Summary: The authors have nothing to disclose.
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