Endoplasmic reticulum chaperone genes encode effectors of long-term memory

Snehayoti Chatterjee1,2, Ethan Bahl3, Utsav Mukherjee1,2,4, Emily N. Walsh1,2,4, Mahesh Shivarama Shetty1,2, Amy L. Yan1,2, Yann Vanrobaeys1,2,3, Joseph D. Lederman2, K. Peter Giese5, Jacob Michaelson6,7,8,9, Ted Abel1,2*

The mechanisms underlying memory loss associated with Alzheimer’s disease and related dementias (ADRD) remain unclear, and no effective treatments exist. Fundamental studies have shown that a set of transcriptional regulatory proteins of the nuclear receptor 4a (Nr4a) family serve as molecular switches for long-term memory. Here, we show that Nr4a proteins regulate the transcription of genes encoding chaperones that localize to the endoplasmic reticulum (ER). These chaperones fold and traffic plasticity-related proteins to the cell surface during long-lasting forms of synaptic plasticity and memory. Dysregulation of Nr4a transcription factors and ER chaperones is linked to ADRD, and overexpressing Nr4a1 or the chaperone Hspa5 ameliorates long-term memory deficits in a tau-based mouse model of ADRD, pointing toward innovative therapeutic approaches for treating memory loss. Our findings establish a unique molecular concept underlying long-term memory and provide insights into the mechanistic basis of cognitive deficits in dementia.

INTRODUCTION

Impaired memory consolidation and the resulting long-term memory loss is an early symptom of Alzheimer’s disease and related dementias (ADRD) (1–3). Memory consolidation requires the transcription of new genes, in sophisticated spatial and temporal patterns, under the control of specific families of transcription factors (4–7). The largest class of transcription regulators in metazoans is composed of the nuclear receptor superfamily (8), which regulates diverse biological processes ranging from metabolism and reproduction to developmental and neuronal function. Among the several subclasses of nuclear receptors, the Nr4a subfamily [Nr4a1 (NUR77, NGF-IB), Nr4a2 (NURR1/HZF-3/NOT/RNR1), and Nr4a3 (NOR1)] has emerged as a critical mediator of long-term memory (4, 7). Notably, these ligand-independent “orphan” receptors are robustly up-regulated in the hippocampus within minutes after learning (3, 4). The learning-dependent expression of the Nr4a genes is regulated by histone acetylation, which is driven by recruitment of adenosine 3′,5′-monophosphate (cAMP) response element–binding protein (CREB)–binding protein (CBP) (9) to CREB response elements in the promoters of these genes (10, 11). Blocking the expression or inactivating the transactivation function of Nr4a factors is sufficient to impair long-term memory (4, 7) and synaptic plasticity (12), whereas transgenic or pharmacological activation enhances long-term memory (13, 14). Moreover, Nr4a function is impaired in brain disorders characterized by debilitating cognitive impairment, ranging from schizophrenia to Parkinson’s disease to ADRD (3, 4, 15, 16). However, despite the critical importance of the Nr4a subfamily, the effector genes that these transcription factors regulate in the hippocampus during memory consolidation have remained elusive. Here, we identify endoplasmic reticulum (ER) chaperone genes as downstream effectors of Nr4a transcription factors, and we establish a role for chaperone function in long-term memory and synaptic plasticity. We further demonstrate that Nr4a transcription factors and the downstream ER chaperones that they regulate are key mediators of the long-term memory loss associated with ADRD, providing candidate targets for the development of innovative therapeutic interventions.

RESULTS

Nr4a transcription factors regulate expression of genes encoding ER chaperones during memory consolidation

To identify effector genes regulated by the Nr4a subfamily during memory consolidation, we used transgenic mice that express a dominant-negative form of Nr4a1 (CaMKIIα-tTA;TetO-Nr4ADN) in excitatory neurons such that the transcriptional activity of all three Nr4a family members is blocked in these cells (4). To assess hippocampus-dependent memory, we examined the performance of Nr4ADN mice in spatial object recognition (SOR), a task that depends on the preference of mice to explore a spatially displaced object (17). In a 24-hour test of long-term memory, the control mice (CaMKIIα-tTA), but not Nr4ADN mice (double transgenic: CaMKIIα-tTA;TetO-Nr4ADN), exhibited preference for the displaced object (Fig. 1A and B). In contrast, in a 1-hour test of short-term memory, both the Nr4ADN and control mice showed a preference for the displaced object (Fig. S1). Thus, Nr4ADN mice exhibit selective deficits in long-term spatial memory. To identify genes regulated by Nr4a transcription factors during memory consolidation, we trained Nr4ADN and control littermates in the SOR task and extracted total RNA from the dorsal hippocampus 2 hours after training (Fig. 1C). We chose this time point to identify effector genes targeted by the Nr4a subfamily of transcription factors, which are immediate early genes induced within minutes after training. This analysis revealed 54 differentially expressed genes (DEGs) (Fig. 1D)
and table S1) in Nr4ADN versus control mice, with 40 down-regulated and 14 up-regulated genes in Nr4ADN mice after learning (Fig. 1D). Enrichment network analysis was used to identify the pathways most represented among the down- and up-regulated genes. The down-regulated pathways included protein processing in the ER, chaperone binding, protein disulfide isomerase activity, and several other pathways related to protein folding in the ER (Fig. 1E). The up-regulated pathways included the poly-pyramidine tract binding pathway linked to Cirbp expression, an RNA binding protein associated with translational control (Fig. 1F). Protein-protein interaction (PPI) analysis of the down-regulated genes in Nr4ADN mice identified a significant cluster composed of nine chaperone proteins (Hspa5, Hsp90b1, Pdia3, Pdia4, Pdia6, Sdf2l1, Dnajb11, Hyou1, and Calr; fig. S2). All of these are components of a large ER multiprotein chaperone complex known to bind nascent proteins (18).

Next, we performed RNA sequencing (RNA-seq) using the dorsal hippocampus from control mice trained in SOR (SOR training +2 hours) or untrained mice [homecage (HC)]. RNA-seq analysis revealed that learning increased the expression of 42 genes and reduced expression of 9 genes (fig. S3 and table S2). Comparison of this gene expression data with the data from control and Nr4ADN mice after learning (Fig. 1D) identified 15 genes induced in control mice after learning that were down-regulated in Nr4ADN mice. These genes included ER chaperone genes Hspa5, Pdia4, Pdia6, Sdf2l1, and Dnajb11 (Fig. 2A). We next analyzed Nr4a1 occupancy on two of these candidate genes that are critical for protein folding (Hspa5 and Pdia6) (18) using data from a previously published Nr4a1 chromatin immunoprecipitation followed by sequencing (ChIP-seq) study (19). The promoters of the Hspa5 and Pdia6 genes were found to be enriched for Nr4a1 binding motifs (fig. S4), suggesting that this
transcription factor directly regulates the expression of these genes. We further examined the expression profiles of these two candidate genes during the first 4 hours after learning in wild-type (C57BL/6J) mice, revealing that spatial learning induced the expression of both Hspa5 and Pdia6 mRNAs in C57BL/6J male mice trained in SOR and euthanized at the indicated times after training (1 hour: n = 9; 2 hours: n = 9; 3 hours: n = 9; 4 hours: n = 9), expressed as fold difference from that of mice handled only in the homecage (HC) (baseline controls, n = 10). One-way ANOVA: Hspa5: F_{4,41} = 13.00, P < 0.0001. Sidak's multiple comparisons tests: ***P < 0.0001 (HC versus 1 hour), ***P < 0.0001 (HC versus 2 hours), **P < 0.001 (HC versus 3 hours), and *P = 0.0158 (HC versus 4 hours); Pdia6: F_{4,41} = 8.442, P < 0.0001. Sidak's multiple comparisons tests: ***P = 0.0008 (HC versus 1 hour), ***P = 0.0001 (HC versus 2 hours), ***P < 0.0001 (HC versus 3 hours), and **P = 0.0054 (HC versus 4 hours). (D and E) Down-regulation of gene expression at 2 hours after SOR training in male Nr4ADN (n = 9) and control (n = 10) mice, as validated by qPCR. Unpaired t test: t_{17} = 3.305, **P = 0.0042 (Hspa5); t_{17} = 3.630, **P = 0.0021 (Pdia6). (F and G) Quantification of Western blot of lysates of synaptosomes isolated from the dorsal hippocampus of male Nr4ADN (n = 4) and control mice 2 hours after SOR training (n = 4). Unpaired t test: t_{6} = 4.011, **P = 0.0070 (Hspa5); t_{6} = 2.982, *P = 0.0246 (Pdia6).

Chaperones, such as Hspa5 and Pdia6, are found in synaptosomes, where they facilitate the folding and assembly of nascent polypeptides, as well as the trafficking of proteins to the neuronal surface (20, 21). The synaptic abundance of Hspa5 and Pdia6 was significantly lower after SOR training in Nr4ADN mice compared to controls (Fig. 2, F and G, and fig. S8). Previous studies have demonstrated that Hspa5 plays a critical role in regulating the postsynaptic membrane delivery of the N-methyl-D-aspartate (NMDA) receptor subunit GluN2A in response to neuronal stimulation (20). Therefore, we next investigated whether Nr4a factors might regulate the surface expression of GluN2A. We expressed Nr4ADN [or enhanced green fluorescent protein (eGFP) as a control] in primary hippocampal neurons using a viral-based approach and assessed the distribution of GluN2A receptors following KCl-mediated neuronal depolarization. We found significant increases in surface levels of GluN2A in eGFP-transduced cells after neuronal depolarization (Fig. 3, A and B), whereas Nr4ADN-transduced cells failed to exhibit activity-induced trafficking of GluN2A (Fig. 3, A and B). We also found that Nr4ADN-expressing neurons show a significant decrease in activity-induced post-synaptic surface localization of GluN2A, as evidenced by the reduced colocalization of GluN2A with the postsynaptic density protein PSD95 (Fig. 3, C and D). Next, we investigated whether rescue of Hspa5 expression would be sufficient to increase post-synaptic localization of GluN2A in Nr4ADN-expressing neurons. Overexpression of Hspa5 increased PSD95 colocalization with GluN2A in Nr4ADN-expressing neurons (Fig. 3, E and F). Our findings demonstrate that the regulation of chaperone protein gene expression, such as Hspa5, by Nr4a transcription factors is
critical for the folding and synaptic trafficking of receptor proteins that are key to synaptic plasticity.

**Restoring protein chaperone function prevents long-term memory and synaptic plasticity deficits in Nr4ADN mice**

Chaperones fold nascent proteins into functional three-dimensional conformations (22), and their up-regulation after learning (Fig. 2) is essential for the activity-dependent processing and trafficking of key synaptic proteins (Fig. 3). Given the roles of chaperones in protein folding, we next performed experiments to determine whether the deficits in memory and synaptic plasticity observed in Nr4ADN mice are related to impairment of this process. First, we examined the effect of phenylbutyrate (PBA), a hydrophobic chemical chaperone, in Nr4ADN mice. PBA interacts with the exposed hydrophobic regions of nascent proteins to facilitate folding (23), partially reverses the mislocalization of proteins (24), and facilitates delivery of proteins that are critical for neuronal plasticity to the cell surface (25–27). PBA has shown promise in rescuing cognitive impairment in several mouse models of neurodegenerative diseases (28–31), and these neuroprotective effects have been attributed to its chaperone activity (28, 32). Notably, systemic delivery of a single dose of PBA to Nr4ADN mice immediately following SOR training rescued long-term memory deficits (Fig. 4A); the same treatment of control mice did not augment long-term memory (Fig. 4B). Because PBA functions as both a molecular chaperone and an inhibitor of histone deacetylases (HDACs), we confirmed that this rescue was not due to changes in expression of Hspa5 and Pdia6 in Nr4ADN mice (fig. S9). In addition, sodium butyrate (NaBu), an HDAC inhibitor that has no molecular chaperone activity (28), failed to rescue memory in Nr4ADN mice (Fig. 4A). This finding is consistent with our previous observation that broad HDAC inhibition is not sufficient to reverse the memory deficits in Nr4ADN mice (4).

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**Fig. 3. Hspa5 regulates activity-dependent surface trafficking of NMDA receptor GluN2A.** (A and B) Cultured neurons were transduced with eGFP or Nr4ADN on DIV 16 or 17 and stimulated with KCl before live staining. (A) Costaining for dendrites (MAP2 antibody) and surface GluN2A by immunofluorescence (IF) in transduced cells after KCl stimulation. Scale bar, 10 μm. (B) Quantification of surface staining for GluN2A in (A) as mean fluorescence intensity. Mixed-effect analysis: significant adeno-associated virus (AAV) construct x treatment interaction: $F_{1,157} = 38.25, P < 0.0001$. Sidak’s multiple comparisons test: ***$P < 0.0001$ (control eGFP versus stimulated eGFP) and ***$P < 0.0001$ (stimulated eGFP versus stimulated Nr4ADN). (C) Costaining for surface GluN2A and PSD95 by IF in transduced cells following KCl stimulation. Scale bar, 10 μm. (D) Quantification of surface GluN2A and PSD95 colocalization in (C). Mixed-effect analysis: significant AAV construct x treatment interaction: $F_{1,69} = 14.57, P = 0.0003$. Sidak’s multiple comparison tests: **$P = 0.0026$ (control eGFP versus stimulated eGFP) and **$P = 0.0012$ (stimulated eGFP versus stimulated Nr4ADN). (E) Cultured neurons were transduced with Nr4ADN or Nr4ADN + Hspa5 on DIV 16 or 17 and stimulated with KCl before live staining. Costaining for surface GluN2A and PSD95 by IF in transduced cells following KCl stimulation. Scale bar, 10 μm. (F) Quantification of surface GluN2A and PSD95 colocalization in (E). Unpaired t test: $t_{30} = 3.727$, ***$P = 0.0008$. 
We previously showed that Nr4ADN mice exhibit deficits in a form of persistent, protein synthesis–dependent long-term potentiation (LTP) induced by repeated spaced high-frequency stimulation of the hippocampal CA1-Schaffer collateral synapses (12). Given that PBA treatment reversed long-term memory deficits in Nr4ADN mice, we examined its effects on this long-lasting form of LTP. Following 20 min of stable baseline field-excitatory postsynaptic potentials (fEPSPs) recordings, hippocampal slices from Nr4ADN mice were treated by bath application of 2 mM PBA [dissolved in the artificial cerebrospinal fluid (aCSF)]. After 40 min of PBA treatment, long-lasting LTP was induced using a spaced four-train stimulation protocol (four 100-Hz, 1-s trains separated by 5 min). Potentiation in slices from Nr4ADN mice decayed quickly, thus showing deficits in the persistence of LTP, as reported in our previous study (12). Treatment with PBA rescued the deficits in long-lasting LTP in Nr4ADN slices leading to persistently enhanced potentiation compared to vehicle-treated slices (PBA-treated: 187.3 ± 16.2%, n = 7 slices, four mice; vehicle-treated: 124.6 ± 14.9%, n = 7 slices, five mice; unpaired t test, *P = 0.0146). Treatment with 2 mM PBA had no significant effect on the baseline responses (pre-drug baseline, 20 min: 100.1 ± 0.13%; post-drug pre-induction baseline, 20 min: 90.63 ± 6.6%; paired t test, *P = 0.2094). The representative fEPSP traces shown are sampled at baseline (black) and at the end of the recording (red). Scale bar (2 mV), 10 ms. Error bars indicate SEM.

Activation of Nr4a1 or ER chaperone function ameliorates memory impairments in an ADRD mouse model

Nr4a transcription factors were previously implicated in Aβ aggregation and memory deficits (33). Therefore, we investigated the expression of Nr4a transcripts in the hippocampus of ADRD patients with increasing grades of pathology. Using a database of RNA-seq and pathological findings from postmortem brain tissue from patients and healthy controls in The Allen Brain Institute Study of Aging, Dementia, and Traumatic Brain Injury (34), we examined the relationship between hippocampal expression of NR4A subfamily members across increasing Cerad (Consortium to Establish a Registry for Alzheimer’s Disease) score, a neuropsychological assessment of the progression of AD, and Braak stage, a measure of the distribution and pathological burden of neurofibrillary tangles (NFTs). We found that levels of expression of NR4A1 and NR4A2 were negatively correlated with both the Cerad scores (Fig. 6A) and Braak stages (Fig. 6B), both measures of the severity of ADRD pathology, whereas NR4A3 was not significantly correlated with disease pathologies. Consistent with our findings, NR4A2 protein expression is reduced in postmortem AD hippocampus at Braak stage VI (33).

Cognitive impairment is a significant feature of ADRD, and we hypothesized that NR4A down-regulation might be related to the compromise in cognitive abilities. To examine this in a mouse model of ADRD, we used the rTg4510 mouse, which overexpresses mutant Hspa5 in the memory deficits observed in Nr4ADN mice, we reinstated Hspa5 expression selectively in hippocampal excitatory neurons using a viral approach and performed behavioral studies 2 weeks after viral infusion (Fig. 5, A to C). The level of overexpression achieved was sufficient to reverse the long-term spatial memory deficits observed in Nr4ADN mice (Fig. 5D), supporting the idea that Hspa5 is downstream of Nr4a transcription factors (Fig. 5E). These findings suggest that the deficits in synaptic plasticity and long-term memory in Nr4ADN mice are due to disruption of a chaperone activity required for the folding of newly synthesized proteins. Overall, we conclude that the activity-induced regulation of ER chaperone Hspa5 by Nr4a transcription factors is essential for the native protein folding required for consolidation of long-term memory.
human tau (tau P301L) exclusively in excitatory neurons (35, 36). These mice develop tangle-like inclusions (35, 36) and show pathological hyperphosphorylation of tau proteins (AT8) in the dorsal hippocampus starting at 3 to 4 months of age (Fig. 6C). These mice have deficits in spatial learning in the Morris water maze, contextual fear conditioning (35–38), and long-term spatial memory in the SOR task (Fig. 6D). Doxycycline treatment prevents these memory deficits (Fig. 6D), demonstrating that it is the expression of the mutant tau transgene and not the transgene insertion site (39) that drives the tauopathy-like phenotype. As in the case of the human postmortem data, the expression of both Nr4a1 and Nr4a2 was down-regulated in the dorsal hippocampus of rTg4510 mice after SOR training (Fig. 6E). These findings validate the appropriateness of using this mouse line as a model for Nr4a dysregulation.

**DISCUSSION**

Here, we show that Nr4a transcription factors act during memory consolidation to drive the expression of genes encoding chaperones that are part of a multiprotein complex within the ER, thereby facilitating folding of the proteins into their functional conformations (18). This study provides functional evidence that these chaperones are involved in synaptic plasticity and long-term memory. The results demonstrating that long-term memory can be reversed in Nr4ADN mice by either application of the chemical chaperone PBA or overexpression of Hspa5 reveal that the protein folding machinery plays a critical role in memory consolidation. Our work in hippocampal neurons identifies the synaptic membrane protein GluN2A as a candidate target protein whose surface trafficking is regulated by Nr4a1-driven expression of Hspa5. Nr4a1 was previously shown to be involved in regulating dendritic spine density (40), consistent with the hypothesis that the target genes of this transcription factor affect synaptic structure and function. Our identification of ER chaperones...
Fig. 6. Down-regulation of Nr4a factors and their effector genes is linked to ADRD pathology. (A and B) Expression profiles of NR4A1, NR4A2, and NR4A3 in the hippocampus, from the Allen Brain Institute Study of Aging, Dementia, and Traumatic Brain Injury, correlated with (A) Cerad scores, which reflect the density of neuritic plaques, and (B) Braak stages, which reflect the severity of neurofibrillary tangles. (C) Schematic depiction of control (CaMKIIα-tTA) or rTg4510 (CaMKIIα-tTA and TetO-hMAPT P301L) mice and Western blots of tau phosphorylation (AT8, phosphorylation at both Ser 202 and Thr 205) in the dorsal hippocampus in the presence or absence of doxycycline (Dox). (D) Long-term memory in rTg4510 and control mice at 4 months of age, following training in SOR. Two-way ANOVA: significant main effect of genotype/treatment ($F_{2,29} = 4.792, P = 0.0159$) and significant main effect of sessions ($F_{1,29} = 9.221, P = 0.0050$). Sidak’s multiple comparisons test: **$P = 0.0096$ (control mice, train versus test), *$P = 0.0439$ (rTg4510-Dox mice, train versus test), **$P = 0.0016$ (rTg4510 mice, 24-hour test versus rTg4510 Dox, 24-hour test), Control, $n = 11$ (3F); rTg4510, $n = 8$ (4F); and rTg4510 + Dox, $n = 13$ (7F). (E and F) rTg4510 and control mice ($n = 9$ per group) were trained in SOR, and 2 hours later, the dorsal hippocampus was collected and processed for RNA extraction and the analysis of gene expression. (E) Nr4a subfamily gene expression: unpaired t test: $Nr4a1$: $t_{16} = 4.878, ***P = 0.0002$; $Nr4a2$: $t_{16} = 2.621, *P = 0.0185$. (F) Hspa5 and Pdia6 gene expression: unpaired t test: $Hspa5$: $t_{16} = 3.692, **P = 0.0020$; $Pdia6$: $t_{16} = 4.177, ***P = 0.0007$. 
The work described here identifies ER chaperone proteins as critical molecular regulators of memory storage. ER chaperones have been studied mainly for their roles in ER stress and the unfolded protein response; their role in memory consolidation is underexplored. Our finding that overexpression of Nr4a1 or Hspa5 chaperone reverses memory deficits in a mouse model of ADRD supports this as an innovative therapeutic approach.

**MATERIALS AND METHODS**

**Data reporting**

No statistical methods were used to predetermine sample size.

**Mouse lines**

**Nr4ADN mice**

Adult Nr4ADN mice were 2 to 4 months old at the time of behavioral or biochemical experiments. They were maintained on a C57BL/6J background and harbored both the CaMKIIα-driven tTA and Tet-O-Nr4ADN transgenes. Incorporation of the CaMKIIα-tTA transgene into chromosome 12 causes a 508.12-kb deletion that affects five genes: Vipr2, Wdr60, D430020J02Rik, Ncapg2, and Ptprn2. To account for any effects of the deleted genes on memory or gene expression, age-matched CaMKIIα-tTA-expressing littermates were used as controls throughout the study.

**rTg4510 mice**

These mice were 3 to 4 months old at the time of behavioral or biochemical experiments. They were maintained on a C57BL/6J background and harbored two transgenes: CaMKIIα-driven TA and Tet-O-driven human tau P301L. Age-matched CaMKIIα-driven TA-expressing littermates were used as controls.

**C57BL/6J mice**

Adult male mice purchased from The Jackson Laboratory were 2 to 4 months of age during behavioral or biochemical experiments. All mice had free access to food and water; lights were maintained on a 12-hour light/12-hour dark cycle.

To suppress TetO-driven expression of Nr4ADN or human tau P301L transgenes, Nr4ADN or rTg4510 mice were placed on a diet...
containing doxycycline (200 mg/kg; Bio-Serv) from weaning until behavioral experiments. All behavioral testing was performed during the light cycle between Zeitgeber time (ZT) 0 to 2. For all behavioral and biochemical experiments, mice were randomly assigned to groups, housed individually for 7 days before experiments, and handled for 2 min per day for 5 days. All experiments were conducted according to U.S. National Institutes of Health (NIH) guidelines for animal care and use and were approved by the Institutional Animal Care and Use Committee of the University of Iowa, Iowa.

**Drugs**

Sodium PBA (Sigma-Aldrich) and NaBu (Sigma-Aldrich) were dissolved in saline. For electrophysiology experiments, the dose of PBA was chosen on the basis of the range of median inhibitory concentration (IC$_{50}$) values reported in the published literature (50). For behavioral experiments, mice were injected with PBA or NaBu intraperitoneally at a dose of 200 mg/kg, immediately after SOR training. Control mice were injected with vehicle (0.9% sterile saline).

**Adeno-associated virus constructs**

AAV$_{2/9}$-CaMKIIα-eGFP, AAV$_{2/9}$-CaMKIIα-Nr4ADN, and AAV$_{2/9}$-CaMKIIα-Hspa5-Tavi, AAV$_{2/2}$-CaMKIIα-Nr4ADN, and AAV$_{2/2}$-CaMKIIα-eGFP were purchased from VectorBuilder (VectorBuilder Inc.).

**Stereotactic surgeries**

Mice were anesthetized using isoflurane and kept on a warm heated pad throughout the stereotactic surgery procedure. Meloxicam was injected as analgesics (51). Viral infusion was performed using a 33-gauge beveled needle [World Precision Instruments (WPI)] attached to a 10-μl Nanofluid syringe controlled by a microsyringe pump (UMP3; WPI). The coordinates for dorsal hippocampus were as follows: anteroposterior, −1.9 mm; mediolateral, ±1.5 mm; and 1.5 mm below bregma. The needle was lowered to the site of injection over 20 min of initial application. In the electrophysiological data presented, n represents the number of slices. Data were acquired using Clampex 10 and Axon Digidata 1440 digitizer (Molecular Devices, Union City, CA) at 20 kHz and were low-pass–filtered at 2 kHz with a four-pole Bessel filter. Data analysis was performed using the Clampfit 10 software (Molecular Devices, Union City, CA). Data were plotted, and statistical analyses were performed using the GraphPad Prism 8 software. For each slice, the fEPSP amplitude was normalized against the average slope over the 20-min baseline (pre-drug baseline in the PBA-treated slices). Data are presented as means ± SEM. LTP persistence was assessed by comparing the final 20-min recordings for the vehicle and treatment groups using a two-way repeated-measures analysis of variance (ANOVA). The mean fEPSP slope of the final 20-min recordings for each group was compared using two-tailed, unpaired t test. Statistical significance was set at P < 0.05.

**Isolation of whole-cell extracts and synaptosomal fractions**

For whole-cell lysate preparation, flash-frozen dorsal hippocampal tissue was homogenized mechanically in 300 μl of ice-cold radioimmuno-precipitation assay (RIPA) buffer (Sigma-Aldrich) supplemented with...
0.2% Triton X-100 (Sigma-Aldrich) and Protease and Phosphatase Inhibitor Cocktail (1:100; Thermo Fisher Scientific). The lysate was kept on ice for 30 min, following which they were centrifuged at 10,000g for 15 min at 4°C. The pellet was discarded, and the supernatant (whole-cell lysate) was collected for Western blot analysis. For synaptosomal extraction, hippocampal tissue was mechanically homogenized in Syn-PER Reagent (Thermo Fisher Scientific) containing Halt Protease and Phosphatase Inhibitor Cocktail (1:100; Thermo Fisher Scientific). The homogenate was centrifuged at 1200g for 10 min at 4°C, after which the pellet (nuclear fraction) was discarded and the supernatant was centrifuged again at 15,000g for 25 min at 4°C. This pellet (synaptosomal fraction) was resuspended in RIPA buffer (Sigma-Aldrich) containing Halt Protease and Phosphatase Inhibitor Cocktail (1:100) and 0.2% Triton X-100 (Sigma-Aldrich).

**Western blot analysis**

Protein extracts were transferred to polyvinylidene difluoride membranes as previously described (3). Membranes were blocked with Odyssey Blocking Buffer in tris-buffered saline (LI-COR) and incubated overnight at 4°C with the following primary antibodies: Hspa5 (1:2000; Proteintech, 11587-1-AP), Pdia6 (1:2000; Abcam, ab11432), Biotin Ligase Epitope Tag (for Tavi-tag detection, 1:1000; Abcam, ab106159), PSD95 (1:5000; Thermo Fisher Scientific, 6G6-1C9), hemagglutinin (HA) (1:2000; Millipore Sigma), phospho-tau AT8 (1:5000; BioLegend, 806503), actin (1:10,000; Thermo Fisher Scientific), ATF4 (1:500; Thermo Fisher Scientific, PA5-27576), ATF6 (1:500; Thermo Fisher Scientific, PA5-114886), and phospho-IRE1α (1:500; Thermo Fisher Scientific, PA5-85738). Membranes were washed and incubated with appropriate IRDye immunoglobulin G (IgG) secondary antibodies, including anti-rabbit IRDye 800LT (1:5000; LI-COR) and anti-mouse IRDye 680CW (LI-COR). Images were acquired using the Odyssey Infrared Imaging System (LI-COR). Quantification of Western blot bands was performed using Image Studio Lite v.5.2 (LI-COR).

**RNA extraction, cDNA synthesis, and quantitative real-time reverse transcription polymerase chain reaction**

Dorsal hippocampi were dissected and immediately stored at −80°C in RNA later solution (Ambion) for later isolation of total RNA. For RNA extraction, Qiagen was added to the hippocampal tissues and they were homogenized using stainless steel beads (Qiagen). Chloroform was then added to the homogenates, and the samples were centrifuged at 12,000g at room temperature for 15 min. RNA was precipitated from the aqueous phase using ethanol and then cleaned using an RNaseasy kit (Qiagen). RNA was eluted in nucleoside-free water and treated with DNase (Qiagen) at room temperature for 25 min. The cleaned RNA was precipitated in ethanol, sodium acetate (pH 5.2), and glycogen overnight at −20°C. RNA samples were centrifuged at top speed at room temperature for 20 min. The precipitates were further washed with 70% ethanol and centrifuged at top speed for 5 min. The RNA precipitates were dried and resuspended in nucleoside-free water, and concentrations were estimated using a NanoDrop (Thermo Fisher Scientific). Complementary DNAs (cDNAs) were prepared from 1 μg of RNA using the SuperScript IV First-Strand Synthesis System (Ambion). Real-time reverse transcription polymerase chain reactions (RT-PCRs) were performed in a 384-well optical reaction plate with optical adhesive covers (Life Technologies). Each reaction was composed of 2.25 μl of cDNA (2 ng/μl), 2.5 μl of Fast SYBR Green Master Mix (Thermo Fisher Scientific), and 0.25 μl of primer mix (IDT). A minimum of three technical replicates per reaction was performed on the QuantStudio 7 Flex Real-Time PCR System (Applied Biosystems, Life Technologies). Data were normalized to housekeeping genes (Tubulin, Pgk1, and B2m), and 2^{−ΔΔCT} method was used for gene expression analysis.

**RNA library preparation, sequencing, and analysis**

RNA libraries were prepared at the Iowa Institute of Human Genetics (IIHG), Genomics Division, using the Illumina TrueSeq Stranded Total RNA with Ribo-Zero gold sample preparation kit (Illumina Inc., San Diego, CA). The KAPA Illumina Library Quantification Kit (KAPA Biosystems, Wilmington, MA) was used to measure library concentrations. Pooled libraries were sequenced on Illumina HiSeq4000 sequencers with 150-bp paired-end chemistry (Illumina) at the IIHG core. RNA-seq data were processed with the bcbio-nextgen pipeline (https://github.com/bcbio/bcbio-nextgen, version 1.1.4). The pipeline uses STAR (54) to align reads to the mm10 genome build (GENCODE release M10, Ensembl 89 annotation) and quantifies expression at the gene level with featureCounts (55). All further analyses were performed using R (56). For gene-level count data, the R package EDASeq (57) was used to adjust for GC content effects (full quantile normalization) and account for sequencing depth (upper quartile normalization) (figs. 10 and 12). Latent sources of variation in expression levels were assessed and accounted for using RUVSeq (RUVr mode using all features) (figs. 11 and 13) (58). Appropriate choice of the RUVSeq parameter k was guided through inspection of P value distributions, relative log expression (RLE) plots, and principal components analysis (PCA) plots. Specifically, the smallest value k was chosen where the P value distribution showed an expected peak below 0.05, RLE plots were evenly distributed and zero-centered, and PCA plots demonstrated replicate sample clustering in the first three principal components (59). Differential expression analysis was conducted using the edgeR quasi-likelihood pipeline (60–62). Codes to reproduce the RNA-seq analysis are available at https://github.com/ethanbahl/chatterjee2021_nr4a.

**Gene Ontology and pathway enrichment analyses of DEGs**

The ClueGO (63) and CluePedia plug-ins of the Cytoscape 3.7.5 software (64) were used in “Functional analysis” mode, using the default parameters for analyzing gene ontology, molecular function, and KEGG pathways in networks for DEGs. The names of significant DEG were pasted into the “Load Marker List” of ClueGO, and the organism “Mus Musculus [10090]” was selected.

**Construction of PPI networks**

The protein-protein interactive network was constructed using STRING (65) (version 11.0), which uses the STRING database (http://string-db.org/) (66). The PPI network was constructed to identify the interactions between proteins encoded by down-regulated DEGs based on experimental data. The DEG names were pasted into “STRING protein query.” Active interaction sources, including text mining, experiments, databases, coexpression, neighborhood, gene fusion, and co-occurrence, were applied, and highest interaction score confidence (0.900) was selected to construct the PPI networks. Full network was constructed, where the edges indicate both functional and physical protein associations.

**ChIP-seq analysis**

Using the SRA toolkit fastq-dump, raw ChIP-seq data from Liu et al. (19) [Gene Expression Omnibus (GEO) accession code GSE96969]
were downloaded, including Nr4a1-HA (SRR6788331), IgG-control (SRR6788333), and input DNA (SRR6788332). The raw reads were trimmed using Trimmomatic version 0.36 with the parameters ILLUMINACLIP:2:30:15 LEADING:30 TRAILING:30 MINLEN:23, and quality inspection was conducted using FastQC version 0.11.5. The trimmed reads from all datasets were aligned using BWA version 0.7.15 with the algorithm mem and default parameters. Signal density files in BedGraph format were generated using BEDTools genomcenv version 2.26.0 with default parameters and then converted in uniform 10-nucleotide bin WIG files for further normalization steps. Peak calling was done using MACS2 version 2.1.2 with a q-value cutoff of 0.01 and default parameters. Each signal density file corresponding to a dataset was scaled such that the total sum of the signal over the mouse mm10 genome was equivalent to 1 million reads of 100 nucleotides. The signal of the input dataset was then subtracted from its corresponding immunoprecipitation dataset to generate the “sciWT-ctrl” files. The normalized WIG files were then encoded in bigWig format using the Kent utilities and visualized using the Integrative Genomics Viewer with the mouse mm10 reference genome.

**Primary neuronal cultures and adeno-associated virus transduction**

Hippocampi from mice (C57BL/6j) were used to generate primary neuronal cultures as previously described (67). Briefly, hippocampi from postnatal P0/P1 pups were dissected, trypsinized (0.25%), and triturated using a fire-polished glass Pasteur pipette to prepare a single-cell suspension. Cells were then plated on poly-l-lysine (1 mg/ml)–coated four-well glass-bottom dishes (Cellvis) at optimal density (150 to 200 cells/mm²) and maintained in Neurobasal medium (Gibco) containing B27 Supplement (Gibco), in an incubator with 5% CO₂ and at 37°C. At days in vitro (DIV) 16 to 17, neurons were transduced with AAV$_{1.2}$-CaMKIIα-Nr4ADN, AAV$_{3.9}$-CamKIIα-Hspa5-Tavi, and AAV$_{2.2}$-CamKIIα-eGFP (titer of concentrated viral stock was $1 \times 10^{13}$ to $2 \times 10^{13}$ genome copies/ml) in a 1:1000 dilution of Neurobasal medium containing B27 Supplement. At 8 to 10 hours following transduction, half of the existing medium was replenished with fresh Neurobasal medium containing B27 Supplement. Cultures were typically maintained until DIV 23 to 25 before experiments commenced.

**KCl stimulation and surface labeling of GluN2A**

At DIV 23 to 25, neurons were incubated in low KCl–Hepes-buffered saline (290 mM osm) [110 mM NaCl, 5.4 mM KCl, 1.8 mM CaCl₂, 0.8 mM MgCl₂, 10 mM d-glucose, and 10 mM Hepes-NaOH (pH 7.4)] for 60 min. Thereafter, neurons were stimulated for another 60 min with high KCl-HBS (same as low KCl-HBS, except for 55 mM NaCl and 60 mM KCl). The high KCl-HBS was washed off, and live neurons were then immunolabeled with N-terminal NMDAR2A antibody (1:25; Thermo Fisher Scientific) in low KCl-HBS for 30 min to exclusively stain the surface GluN2A receptors. Following the antibody incubation, the cells were washed twice with phosphate-buffered saline containing Mg$^{2+}$ and Ca$^{2+}$ (PBS-MG; 137 mM NaCl, 2.7 mM KCl, 10 mM Na$_2$HPO$_4$, 2 mM KH$_2$PO$_4$, 1 mM MgCl$_2$, and 0.1 mM CaCl$_2$). Cells were then fixed in PBS-MG containing 2% paraformaldehyde and 2% sucrose for 15 min at 37°C, washed thrice in PBS-MG at room temperature, and blocked with PBS-MG containing 2% bovine serum albumin for 60 min at room temperature. Cells were incubated with Alexa Fluor 647–conjugated goat anti-rabbit secondary antibody (1:200; Invitrogen) at room temperature for 90 min in blocking solution. Neurons were then permeabilized with PBS-MC containing 0.1% Triton X-100 at room temperature for 5 min and incubated with blocking solution for 60 min and thereafter with MAP2 antibody (1:1000; Sigma-Aldrich) for 8 to 10 hours at 4°C. Cells were washed thrice in PBS-MC and incubated with Alexa Fluor 546–conjugated goat anti-mouse secondary antibody (1:200; Invitrogen) at room temperature for 60 min. Last, cells were washed three times with PBS-MC at room temperature and preserved in PBS-MC for future imaging.

**Confocal imaging and image analysis**

Cultured neurons after completion of the experiments were imaged using an Olympus FV3000 confocal microscope with a 100x numerical aperture $= 1.45$ oil immersion objective at 1024 x 1024 pixel resolution. High-magnification images were captured using 3x optical zoom. All images (8 bit) were acquired with identical settings for laser power, detector gain, and pinhole diameter for each experiment and between experiments. Images from the different channels were stacked and projected at maximum intensity using ImageJ (NIH). Mean fluorescence intensity of surface GluN2A and the colocalization between surface GluN2A and PSD95 puncta were assessed using plugins in ImageJ.

**Analysis of human AD**

Using RNA sequencing data from the “Aging, Dementia, and TBI Study” (34) provided by the Allen Institute for Brain Science, we fit linear models between the RNA integrity number (RIN)–corrected log$_2$-transformed hippocampal expression levels of NR4A family members and the individual’s Cerad score, a semiquantitative estimate of neuritic plaque density (34).

**Statistics**

Behavioral and biochemical data were analyzed using paired or unpaired two-tailed $t$ tests and either one-way or two-way ANOVAs (in some cases, with repeated measures as the within-subject variable). Sidak’s tests were used for post hoc analyses, where needed. Differences were considered statistically significant when $P < 0.05$. As indicated for each figure panel, all data are plotted in either bar graphs, in which symbols represent each data point, or dot plots, where each symbol represents an individual data point. Graphs were plotted as means ± SEM.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abm6063

View/request a protocol for this paper from Bio-protocol.

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