Small molecule cognitive enhancer reverses age-related memory decline in mice.

SHORT TITLE (less than 40 characters): Reversing aging related deficits

Authors: Karen Krukowski1,2, Amber Nolan2,3,*, Elma S. Frias1,2,*, Morgane Boone4,*. Gonzalo Uretas5, Katherine Grue1,2, Maria-Serena Paladini1,2, Edward Elizarraras1,2, Luz Delgados, Sebastian Bernales5, Peter Walter4,6 and Susanna Rosi1,2,7,8,9

1 Department of Physical Therapy and Rehabilitation Science, University of California at San Francisco, San Francisco, CA, USA.
2 Brain and Spinal Injury Center, University of California at San Francisco, San Francisco, CA, USA.
3 Department of Pathology, University of California at San Francisco, San Francisco, CA, USA.
4 Biochemistry and Biophysics, University of California at San Francisco, San Francisco CA, USA.
5 Fundación Ciencia & Vida, Santiago, Chile.
6 Howard Hughes Medical Institute, University of California at San Francisco, San Francisco, CA, USA.
7 Department of Neurological Surgery, University of California at San Francisco, San Francisco, CA, USA.
8 Weill Institute for Neuroscience, University of California at San Francisco, San Francisco, CA, USA.
9 Kavli Institute of Fundamental Neuroscience, University of California at San Francisco, San Francisco, CA, USA.

*authors contributed equally

Corresponding Authors:

Susanna Rosi, Ph.D. Lewis and Ruth Cozen Chair II
Professor
1001 Potrero Ave,
Zuckerberg San Francisco General Hospital
Building #1 Room 101
San Francisco, CA 94110
Tel.: +1-415-206-3708
Email: susanna.rosi@ucsf.edu

Peter Walter, PhD, Professor
Box 2200, UCSF
San Francisco, CA 94143
Phone: +1-415-476-5017
Email: peter@walterlab.ucsf.edu
ONE SENTENCE SUMMARY
Inhibition of the integrated stress response restores neuronal and immune dysfunction and alleviates memory deficits in aged mice.
ABSTRACT

With increased life expectancy, age-associated cognitive decline becomes a growing concern. The integrated stress response (ISR) is activated during aging and contributes to age-related brain phenotypes. We demonstrate that treatment with the drug-like small-molecule ISR inhibitor ISRIB reverses ISR activation in the brain, as indicated by decreased activating transcription factor 4 (ATF4) protein levels. Furthermore, ISRIB treatment reverses spatial memory deficits and ameliorates working memory in old mice. At the cellular level in the hippocampus, ISR inhibition i) rescues intrinsic neuronal electrophysiological properties, ii) restores spine density and iii) reduces immune profiles, specifically interferon and T cell-mediated responses. Thus, pharmacological interference with the ISR emerges as a promising intervention strategy for combating age-related cognitive decline.

INTRODUCTION

"Of the capacities that people hope will remain intact as they get older, perhaps the most treasured is to stay mentally sharp" (1).

The impact of age on cognitive performance represents an important quality-of-life and societal concern, especially given our prolonged life expectancy. Decreases in executive function as well as learning and memory decrements in older individuals are common (2, 3, 4, 5). According to the US Department of Commerce the aging population is estimated by 2050 to reach 83.7 million individuals above 65 years of age in the US; this represents a rapidly growing healthcare and economic concern (6).
Age-related decline in spatial memory has been recapitulated in preclinical studies with old rodents (7-10). The hippocampus is the brain region associated with spatial learning and memory formation and is particularly vulnerable to age-related changes in humans and rodents (11-14). Deficits in a number of cellular processes have been suggested as underlying causes based on correlative evidence, including protein synthesis (15), metabolism (16), inflammation (17), and immune responses (9, 18-20). While providing a wealth of parameters to assess, by and large the causal molecular underpinnings of age-related memory decline have remained unclear.

The principle that blocking protein synthesis prevents long-term memory storage was discovered many years ago (21). With age there is a marked decline of protein synthesis in the brain that correlates with defects in proper protein folding (22-24). Accumulation of misfolded proteins can activate the integrated stress response (ISR) (25), an evolutionary conserved pathway that decreases protein synthesis. In this way, the ISR may have a causative role in age-related cognitive decline. We previously discovered that interference with the drug-like small-molecule inhibitor (integrated stress response inhibitor, or ISRIB) rescued traumatic brain injury-induced behavioral and cognitive deficits (26, 27), suggesting that this pharmacological tool may be useful in testing this notion.

Increasing age leads to structural and functional changes in hippocampal neurons. Specifically, in old animals there is an increase in neuronal hyperpolarization after spiking activity (“afterhyperpolarization”, or AHP) that decreases intrinsic neuronal excitability and...
correlates with memory deficits \((11-14, 28)\). Aging also manifests itself with synaptic excitability changes in the hippocampus that correlate with a reduction in the bulbous membrane projections that form the postsynaptic specializations of excitatory synapses, termed dendritic spines \((29, 30)\). Morphological changes in dendritic spine density is critical for spatial learning and memory \((31, 32)\). Whether these age-related neuronal changes can be modified or are linked with ISR activation has yet to be determined.

In addition to neuronal changes, ISR activation can modify immune responses via alterations in cytokine production \((33)\). Indeed, maladaptive immune responses have been linked with cognitive decline in the old brain \((8, 9, 18, 20)\). Initial studies focused on age-associated cytokine responses, including interferon (IFN)-mediated cognitive changes \((18, 34)\). Type-I IFN responses can induce age-related phenotypes in rodents. Furthermore, the adaptive immune system (T cell infiltration into the old brain) can regulate neuronal function via IFN-γ production \((19)\), suggesting the possibility that age-induced maladaptive immune responses and the ISR are linked. Here we explore the possibility of ISR inhibition by ISRIB as a potential strategy for modifying age-induced neuronal, immune, and cognitive dysfunction.

RESULTS

ISRIB resets the ISR in the brain of old mice.
ISR activation leads to global reduction in protein synthesis but also to translational up-regulation of a select subset of mRNAs whose translation is controlled by small upstream open-reading frames in their 5’-UTRs (35, 36). One well-studied ISR-upregulated target protein is ATF4 (activating transcription factor 4) (37, 38). We recently showed ISRIB administration reversed mild head trauma-induced elevation in ATF4 protein (27). Using the same ISRIB treatment paradigm of 3 injections on consecutive days (26, 27), we found decreased age-associated ATF4 protein levels in mouse brain lysates when compared to vehicle-treated controls during ISRIB administration (Figure 1A, B; Supplemental Figure 1). Similar trends were observed in lysates from hippocampal tissue (Supplemental Figure 2). ATF4 levels 18 days after cessation of ISRIB treatment showed persistent reduction in age-induced ATF4 protein levels that were indistinguishable from young mice (Figure 1A, C). Thus, acute ISRIB administration in the old brain has long-lasting, chronic effects on ISR activation.

Inhibition of the ISR reverses age-induced decline in spatial learning and memory.

To assess whether the reduction in ISR activation affects age-related cognitive defects, we tested the capacity for spatial learning and memory in young and old mice in a radial arm water maze (26, 39). In this forced-swim behavioral test, animals were trained for two days (two learning blocks/day) to locate a platform hidden under opaque water in one of the eight arms using navigational cues set in the room (Figure 2A). We recorded the total number of entries into the non-target arm (errors) before the animal found the escape
platform with automated tracking software and used it as a metric of learning. After two days of training, young animals averaged one error prior to successfully locating the escape platform, while old animals averaged three errors, indicating their reduced learning capacity (Supplemental Figure 3A).

We next tested whether pharmacological inhibition of the ISR could modify the age-related spatial learning deficits. ISRIB treatment started the day before the first training day and continued with daily injections throughout the duration of the training (3 injections in total; see Figure 2A, left). By the end of two days of training, ISRIB-treated old animals averaged two errors prior to finding the escape platform, while vehicle-treated old animals averaged three, denoting significant learning improvement in the mice that received ISRIB (Supplemental Figure 3B). No difference in learning performance was measured in young mice that received the identical treatment paradigm (Supplemental Figure 3C), suggesting that ISRIB-induced learning improvement measured in this training regime is age-dependent. These results were confirmed in an independent old animal cohort, in which we tested an additional ISR inhibitor (Cmp-003, a small molecule with improved solubility and pharmacological properties (PCT/US18/65555)), using an identical training/injection paradigm (Supplemental Figure 3D). Old animals that received Cmp-003 made significantly fewer errors prior to locating the escape platform than old animals that received vehicle injections, again indicating significant learning improvement.

Spatial memory of the escape location was measured eight days later by reintroducing the animals into the pool and measuring the number of errors before they located the
hidden platform. The animals did not receive any additional treatment during this task. Old mice treated with ISRIB one week before made significantly fewer errors compared to matched, vehicle-treated old male (Figure 2B) and female (Supplemental Figure 4) mice. Remarkably, the memory performance of old animals treated with ISRIB a week before was comparable to that of young mice (Figure 2B). These results demonstrate that acute treatment with ISRIB in old mice rescues chronic age-induced spatial learning deficits, cementing a causative role of the ISR in rendering long-term memory consolidation dysfunctional.

ISRIB administration improves age-induced deficits in working and episodic memory weeks after treatment.

Given the long-lasting effect of acute ISRIB treatment on ATF4 protein levels in the brain and on memory function one week after drug administration, we next tested the duration of ISRIB effects on age-related cognitive function. On day 20 post ISRIB treatment (Figure 2A, right), we measured working and episodic memory using a delayed-matching-to-place paradigm (DMP) (26, 40) in the same animal cohort without additional ISRIB treatment. During DMP animals learned to locate an escape tunnel attached to one of 40 holes in a circular table using visual cues. The escape location was changed daily, forcing the animal to relearn its location. To quantify performance, we used analysis tracking software to measure “escape latency”, reporting the time taken by the mouse to enter the escape tunnel.
Old mice that received ISRIB treatment 20 days earlier displayed significant improvement over the four-day testing period (Figure 2C; Day 20 vs. Day 23). By Day 23 post-treatment animals were locating the escape tunnel on average 20 seconds faster than the matched-vehicle group (Figure 2C). This behavior is indicative of improved working and episodic memory. By contrast, old animals that received vehicle injections did not learn the task (Figure 2C, D; Day 20 vs. Day 23), as previously observed (8, 10). The performance improvement, measured as escape time on Day 20 minus escape time on Day 23, was significantly elevated in ISRIB-treated when compared to the vehicle-treated group (Figure 2D). These results demonstrate that ISRIB administration increases cognitive performance in a behavioral paradigm measured weeks after administration.

**ISRIB treatment reverses age-associated changes in hippocampal neuron function.**

To determine how ISRIB treatment might improve cognition, we investigated the effects of ISRIB treatment on hippocampal neuronal function. Utilizing whole cell patch clamp, we recorded neuronal electrophysiological properties, including the afterhyperpolarization (AHP), in hippocampal CA1 pyramidal neurons of young mice and old mice with vehicle injections, and old mice subjected to a single injection of ISRIB (Figure 3A).

To evaluate for alternations in intrinsic neuronal excitability, we measured action potentials and membrane properties in response to a series of hyperpolarizing and
depolarizing current steps (20 steps from -250 to 700 pA, 250 msec duration), including
the AHP following ~50 Hz spiking activity (i.e. post-burst AHP) induced with a current
step (Figure 3A). In agreement with previous reports (11-14, 28), old mice displayed a
significantly increased AHP amplitude when compared to young mice (Figure 3B). ISRIB
treatment reversed the age-induced increase in AHP amplitude, rendering ISRIB-treated
old mice indistinguishable from young ones (Figure 3B). We measured no significant
differences in other passive membrane and active spiking properties between groups
(Supplemental Figure 5). In addition, spontaneous excitatory postsynaptic currents
(sEPSC) were indistinguishable between groups (Supplemental Figure 6). These data
suggest that ISRIB treatment in old animals restores intrinsic excitability by specifically
reducing the post-burst AHP back to a youthful level.

ISRIB treatment reduces dendritic spine loss.

To determine if ISRIB might affect age-induced synaptic structural changes, we quantified
dendritic spine density after ISRIB treatment in old mice with fluorescently labeled
excitatory neurons (marked by a genomically encoded Thy1-YFP fusion protein). The
hippocampus of old mice is characterized by a reduction in dendritic spine numbers that
correlates with diminished cognitive output (29, 30). Old Thy1-YFP expressing mice
received ISRIB treatment and two days of behavioral training as described in Figure 2A.
At the end of Day 2, we terminated the animals and harvested the brains for quantification
of dendritic spine numbers in the hippocampus (stratum radiatum of CA1) (Figure 3C)
using confocal microscopy imaging and unbiased analysis (Figure 3D). Similar to previous reports, we measured a significant reduction in dendritic spine density in old mice when compared to young Thy1-YFP mice (Figure 3E) (29, 30). ISRIB treatment significantly increased spine numbers when compared with age-matched vehicle-treated mice (Figure 3E). Taken together these data demonstrate that within 1-2 days of ISRIB administration both neuron structure and function in old mice are restored to those of young animals.

**Age-induced interferon (IFN) and T cell responses are reduced following ISRIB treatment.**

To determine differential gene expression patterns impacted by ISRIB administration in the old brain, we performed an unbiased bulk RNA sequencing analysis of hippocampus extracts from three old vehicle-treated and old ISRIB-treated mice after two ISRIB administrations. Due to the heterogeneity observed in old animals, our initial analysis did not reveal significant differences between groups (Supplemental Figure 7A). However, we observed strong trends when a single old outlier mouse was removed. A number of immune and immune regulatory pathways (118 genes total), in particular those related to IFN signaling, were down-regulated after ISRIB treatment (Figure 4A and Supplemental Figure 7). Since immune dysregulation in the brain increases with age (41) (Supplemental Figure 8) and is known to correlate with reduced cognitive performance in the old animals (8, 9, 18, 20), we decided to follow up on these statistically unsubstantiated yet intuitively appealing clues. To this end, we next performed quantitative PCR (qPCR) analyses on hippocampi from young, old, and old + ISRIB.
animals immediately after two ISRIB administrations. These analyses confirmed that age
increased expression of a number of IFN response pathway genes, Ifit1, Gbp10, and Rtp4
(Figure 4B-D). Importantly, ISRIB administration reduced expression of Ifit1, Gbp10, and
Rtp4 to levels indistinguishable from young animals (Figure 4B-D).

We next measured T cell responses in old mice with or without ISRIB treatment (two
injections, Figure 2A). Similar to other reports (19, 42), we observed a significant
increase in T cell marker expression (CD3) in the hippocampus of old compared to young
mice (Figure 4E). ISRIB treatment in the old mice reduced the expression of the T cell
marker comparable to that observed in young mice (Figure 4E), suggesting a link
between the IFN response and T cell recruitment to the brain. T cell marker expression
in the brain correlated with cognitive performance: mice with lower T cell marker
expression made fewer errors prior to locating the escape platform during memory testing
on Day 2 (Figure 4F). The ISRIB-induced reduction in T cell marker levels was not limited
to the brain but extended to the peripheral blood of old animals, with CD8+ T cell
percentages reduced following ISRIB administration (Figure 4G). By contrast, we
observed no changes in CD4+ T cell levels (Figure 4H). Overall, ISRIB treatment impacts
immune parameters both in the periphery and in the brain.

CONCLUSION

We provide evidence for a direct involvement of the ISR in age-related cognitive decline.
Temporary treatment with ISRIB causes down-regulation of ATF4 that is sustained for at
least 20 days. This “ISR reset” leads to improvement in spatial and working memory. At
a cellular level the cognitive enhancement is paralleled by i) improved intrinsic neuron
excitability, ii) increased dendritic spine density and iii) reversal of age-induced changes
in IFN and T cell responses in the hippocampus and blood. Thus, we identify broad-
spectrum anatomical, cellular, and functional changes caused by ISR activation in old
animals. If these findings in mice translate into human physiology, they offer hope and a
tangible strategy to sustain cognitive ability as we age.
FIGURE LEGENDS

Figure 1. ISRIB down-regulates the ISR in the brain of old mice. (A) Experimental dosing scheme- ISRIB treatment denoted by syringes (3 injections). ISRIB treatment reduced ATF4 protein levels (B) acutely during drug administration and (C) chronically 18 days after ISRIB treatment was complete. ATF4 protein levels normalized to actin and graphed relative to old mice. (B) Old males (7) and females (3): Vehicle n = 10; ISRIB n = 10. Student’s t-test. *p < 0.05. (C) Young and old males. Young n = 5, Old = 3, Old + ISRIB = 3. One-way ANOVA (F = 18.8, p < 0.001); with Tukey post-hoc analysis. *p < 0.05; **p < 0.001. Individual animal values represented by dots; lines depict group mean ± SEM.

Figure 2. Inhibition of the ISR reverses age-induced decline in spatial learning and memory. (A) Experimental Design: Old (~19 months) animals underwent behavioral analysis in a radial arm water maze (RAWM) and a delayed matching to place paradigm (DMP). ISRIB or vehicle administration (2.5 mg/kg intraperitoneal) occurred daily during the learning phase of RAWM denoted by syringes (days 0-2). (B) ISRIB treatment improved memory one week after administration in male rodents. One-way ANOVA (F = 5.3, p < 0.05); with Tukey post-hoc analysis. Young n = 10; Old n = 35; Old + ISRIB n = 21. (C,D) Age-induced deficits in working and episodic learning and memory restored weeks after ISRIB administration. Animals performed the DMP from day 20 - 23. (C) Average of all trials per group for each day. Two-way repeated measures ANOVA reveals a significant difference between groups p < 0.01 (denoted in figure legend) and time effect p < 0.01. (D) ISRIB animals displayed significant improvement when compared with the vehicle group. Student t-test. *p < 0.05, **p < 0.01. Old n = 18; Old + ISRIB n = 16. Individual animal scores represented by dots; lines depict group mean ± SEM.

Figure 3. ISRIB treatment alleviates age-associated changes in CA1 pyramidal neuron function and structure. (A) Left: Image of pipette patched onto CA1 neuron in sagittal slice of hippocampus. Right: Representative traces from hippocampal CA1 pyramidal neurons from old animals treated with either vehicle (light blue) or ISRIB (dark blue) or young animals treated with vehicle (orange) showing the response to a current injection eliciting >50Hz spiking activity. Spikes are truncated (dashed line), and the AHP is visualized immediately following cessation of current injection (yellow square) and quantified as the change in voltage from baseline (dotted line). (B) Age-induced increases in AHP were measured when comparing young and old animals. ISRIB treatment reversed increased AHP to levels indistinguishable from young animals. One-way ANOVA (F = 4.461, p < 0.05); with Tukey post-hoc analysis. *p < 0.05. Each neuron is represented with a symbol; lines indicate the mean ± SEM (Neurons: n = 10 young + vehicle (5 animals); n = 12 old + vehicle (5 animals), n = 19 old + ISRIB (7 animals) with 1-5 neurons recorded per animal. (C-E) Spine density was quantified in the CA1 region of the dorsal hippocampus from young and old Thy1-YFP-H mice. (C) Diagram of hippocampal region analyzed. SR = stratum radiatum. (D) Representative images from Old and Old + ISRIB mice. (E) A decrease in dendritic spine density was measured when comparing old mice to young mice. ISRIB treatment significantly increased spine density
levels of old mice when compared to vehicle treated old mice. 63x magnification with a water immersion objective. Young males n = 7 slides (2 animals); Old males + Vehicle n = 12 slides (3 mice); Old males + ISRIB n = 17 slides (4 mice). Individual slide scores (relative to old mice) represented in dots, lines depict group mean ± SEM. One-way ANOVA (F = 18.57, p < 0.001) with Tukey post-hoc analysis. **p < 0.01; ***p < 0.001.

Figure 4. Age-induced IFN and T cell responses are reduced following ISRIB treatment. (A) Bulk RNA-sequencing analysis revealed that of the 129 differentially expressed genes (red dots, FDR < 0.05 and min. 1.2-fold change), 118 genes were down regulated with ISRIB treatment, with a strong enrichment for IFN pathway genes (red dots with black line). DE = differentially expressed. (B-D) IFN response pathways (Ilf1, Gbp10, Rtp4,) were investigated in the hippocampus of young and old mice by qPCR analysis. ISRIB administration reversed age-induced increases in (B) Ilf1, One-way ANOVA (F = 8.8; p < 0.01) with a Tukey-post analysis. Young n = 8; Old n = 7; Old + ISRIB n = 8) (C) Gbp10, One-way ANOVA (F = 4.2, p < 0.05) with a Tukey-post analysis. Young n = 8; Old n = 7; Old + ISRIB n = 7) and (D) Rtp4, One-way ANOVA (F = 12.23, p < 0.001) with a Tukey-post analysis. Young n = 8; Old n = 7; Old + ISRIB n = 8) levels. *p < 0.05; **p < 0.01; ***p < 0.001. (E) CD3 gene-expression changes in the hippocampus of young and old animals were measured by qPCR analysis, comparing expression levels between young, old and old + ISRIB. CD3, a marker for T cells, was significantly increased with age. ISRIB administration returned CD3 expression levels to those comparable to young animals. One-way ANOVA (F = 5.2; p < 0.05). Tukey-post hoc analysis. Young n = 8; Old n = 7; Old + ISRIB n = 8. (F) A significant positive correlation was measured between T cell expression levels in the hippocampus and cognitive performance. Linear regression was measured by Pearson R correlation (R² = 0.27; F = 8.0, p < 0.001). (G, H) Peripheral T cell levels were measured by flow cytometric analysis of whole blood. (G) ISRIB treatment reduced CD8+ T cell percentages (of CD45+ cells) in the peripheral. Student t-test. Old n = 7; Old + ISRIB n = 8. *p < 0.05. (H) CD4+ T cell percentages (of CD45+ cells) were not impacted. Individual animal scores represented by dots; lines depict group mean ± SEM.

Supplemental Figure 1. ISRIB down-regulates ISR-induced pathways. The impact of ISRIB on known ISR activation pathways was investigated by Western blot analysis of brain lysates (raw Western blot data). Each lane represents an individual animal brain extract. (A) Acute end points. (B) Chronic end points.

Supplemental Figure 2. Impact of ISRIB on ISR-induced pathways in the hippocampus. The impact of ISRIB on known ISR activation pathways was investigated by Western blot analysis of hippocampal lysates (A). Raw Western blot data. Each lane represents an individual animal hippocampus extract. (B) ISRIB treatment reduced ATF4. ATF4 protein levels normalized to actin graphed relative to old mice: Old n = 10; Old + ISRIB n = 10. Student’s unpaired t-test. Individual animal scores represented by dots; lines depict group mean ± SEM.
Supplemental Figure 3. ISR inhibitors relieve age-induced deficits in spatial learning. RAWM was used to measure age-induced deficits in spatial learning. Animals ran 2 blocks on each learning day. (A) Old animals performed significantly worse than young animals. Two-way repeated measures ANOVA revealed a significant interaction (p < 0.001). Bonferroni post-hoc to determine differences at various blocks. Old males n = 19, Young males n = 10. *p < 0.05. (B) ISRIB or vehicle administration (2.5 mg/kg intraperitoneal) occurred days 0-2. Compared with the old group, ISRIB treated animals made significantly fewer errors over the course of learning. Two-way repeated measures ANOVA reveals a significant difference between groups p < 0.05. Old n = 19; Old + ISRIB n = 15. (C) No differences were measured between young +/- ISRIB administration. Two-way repeated measures ANOVA revealed no significant differences. Young males + vehicle n = 10; Young males + ISRIB n = 10 (D) Cmp-003 (5 mg/kg intraperitoneal) administration occurred days 0-2. Old mice that received Cmp-003 performed significantly better than old mice that received vehicle. Two-way repeated measures ANOVA revealed a significant group (p < 0.01) and time effect (p < 0.05). Old males n = 9, Old males + Cmp-003 n = 9. **p < 0.01. Data are means ± SEM.

Supplemental Figure 4. ISRIB reduces age-induced memory deficits in female mice. RAWM was used to measure age-induced deficits in learning and memory. ISRIB treatment improved memory one week after administration in female rodents. Two-way ANOVA reveals a significant sex (p < 0.05) and treatment effect (p < 0.01). Old female n = 12; Old female + ISRIB n = 11. ** p < 0.01. Individual animal scores represented by dots, lines depict group mean and SEM.

Supplementary Figure 5. Age and ISRIB treatment do not modify other passive or active intrinsic membrane properties in CA1 pyramidal neurons. (A) Representative traces from CA1 pyramidal neurons showing the membrane potential response to a 250 pA current injection in neurons from old animals treated with either vehicle (light blue) or ISRIB (dark blue) or young animals treated with vehicle (orange). Quantification of the action potential (AP) including the half width (B), amplitude (C), and threshold (D) did not show significant differences between CA1 pyramidal recordings from old, old + ISRIB-treated, or young mice. Likewise, evaluation of the maximum firing frequency (E) or how the frequency of spiking changes over time, quantified by the adaptation index (F) or with current injection, quantified by the slope of the relationship of firing frequency versus amplitude of current injection (F/I slope) (G) was also not significantly different between groups. Finally, passive membrane properties including the membrane time constant (tau) (H), membrane resistance (Rm) (I), and the resting membrane potential (J) were not significantly altered by age or ISRIB treatment. Each neuron is represented with a symbol; solid lines indicate the mean ± SEM. (One-way ANOVA for all comparisons; Neurons: and n = 12 young + vehicle (5 animals); n = 15 old + vehicle (5 animals), n = 22 old + ISRIB (7 animals) with 1-5 neurons recorded per animal.

Supplemental Figure 6. Age and ISRIB treatment do not affect spontaneous excitatory post-synaptic currents (sEPSC) in CA1 pyramidal neurons. (A) Representative whole cell voltage-clamp recordings showing sEPSCs from CA1 pyramidal neurons from old animals treated with either vehicle (light blue) or ISRIB (dark...
blue) or young animals treated with vehicle (orange). Arrows denote synaptic currents.

(B) The sEPSC amplitude was not significantly different between groups (one-way ANOVA). (C) The sEPSC frequency was unchanged after ISRIB treatment or compared to young mice (Kruskal-Wallis test). The median amplitude or frequency for each neuron is represented with a symbol; solid lines indicate the mean ± SEM. (Neurons: n = 11 young + vehicle (5 animals); n = 15 old + vehicle (5 animals), n = 18 old + ISRIB (7 animals) with 1-5 neurons recorded per animal.)

Supplemental Figure 7. Impact of age and ISRIB on myeloid and inflammatory gene signatures. (A) Multidimensional scaling analysis of the hippocampal transcriptome of 3 ‘old’ and 3 ‘old + ISRIB-treated’ mice to visually represent the similarity of the RNA-seq data between mice. Including all mice in the analysis did not reveal a clear separation of groups, as mouse 3 of the ‘old’ cohort clusters together with the ‘old + ISRIB-treated’ mice, suggesting a similar transcriptomic profile. However, ‘old’ mouse 1 and 2 clearly cluster separately from the ‘old + ISRIB-treated’ mice, hinting at a clear transcriptional difference between both groups. (B-D) Transcriptional changes in the hippocampus of old mice point to a downregulation of innate immune and interferon signaling pathways in the brain upon treatment with ISRIB. Pathway enrichment of the 129 differentially expressed genes (DEGs) was calculated across the default databases using the DAVID Functional Annotation Clustering tool (B) and against the Gene Ontology (GO) database (C). DAVID functional clusters with Normalized Enrichment Score (NES) > 2 were manually annotated with the most appropriate overlapping term, while for GO enrichment, the top ten strongest hits are visualized. In parallel, Gene Set Enrichment Analysis across the entire ranked transcriptome also confirmed significant (FDR < 0.05) enrichment of the hallmark IFNγ and IFNα pathways from the MSigDB database (D). Gene scores were calculated as sign(log2FC) * (-log(adj.p-value)). Vertical black lines delineate the genes associated with the given pathway and their gene score rank in the transcriptional data along the x-axis. A negative enrichment score (green line) indicates enrichment of the pathway in downregulated genes. (E) 74.4% (96/129) of genes that change with ISRIB are interferon-responsive genes (as annotated in the Interferome v2.01 database, min. 2-fold change in mouse and human brain-specific datasets), with most being common targets of both type I (incl. IFNα, IFNβ) and type II (IFNγ) IFN responses. (F) Transcription factor (TF) motifs for Interferon Response Factors (IRF) are the top hit when scanning the first 500 bp upstream of the transcriptional start site of DEGs, indicating that with ISRIB, the transcriptional changes are driven by interferon response factors. Motifs and associated TFs are clustered according to motif similarity with the Cytoscape iRegulon plugin.

Supplemental Figure 8. Impact of age on inflammatory expression profiles. (A-D) Gene-expression changes in the hippocampus of young and old animals were measured by qPCR analysis. CCL2, a chemotactic ligand that recruits myeloid and some lymphoid cells into the brain parenchyma, was significantly increased with age. Pro-inflammatory cytokines IL-6, IL-1β, TNFα were modestly increased when comparing age and young animals. Student t-test *p < 0.05. (A-C) Old n = 8, Old + ISRIB n = 7. (D) Old n = 7, Old + ISRIB n = 7. Individual animal values represented by dots; lines depict group mean and SEM.
METHODS

Animals. All experiments were conducted in accordance with National Institutes of Health (NIH) Guide for the Care and Use of Laboratory Animals and approved by the Institutional Animal Care and Use Committee of the University of California, San Francisco (Protocol 170302). Male and female C57B6/J wild-type (WT) mice were received from the National Institute of Aging. Thy-1-YFP-H (in C57 background) were bred and aged in house. Old animals started experimentation at ~19 months of age and young animals 3-6 months of age. Animal shipments were received at least one week prior to start of experimentation to allow animals to habituation the new surroundings. Mice were group housed in environmentally controlled conditions with reverse light cycle (12:12 h light:dark cycle at 21 ± 1 °C; ~50% humidity) and provided food and water ad libitum. Behavioral analysis was performed during the dark cycle.

Drug Administration. ISRIB solution was made by dissolving 5 mg ISRIB in 2.5 mLs dimethyl sulfoxide (DMSO) (PanReac AppliChem, 191954.1611). The solution was gently heated in a 40 °C water bath and vortexed every 30 s until the solution became clear. Next 1 mL of Tween 80 (Sigma Aldrich, P8074) was added, the solution was gently heated in a 40 °C water bath and vortexed every 30 s until the solution became clear. Next, 10 mL of polyethylene glycol 400 (PEG400) (PanReac AppliChem, 142436.1611) solution was added gently heated in a 40 °C water bath and vortexed every 30 s until the solution became clear. Finally, 36.5 mL of 5% dextrose (Hospira, RL-3040) was added. The solution was kept at room temperature throughout the experiment. Each solution was used for injections up to 7 day maximum. The vehicle solution consisted of the same chemical composition and concentration (DMSO, Tween 80, PEG400 and 5% dextrose). Stock ISRIB solution was at 0.1 mg/ml and injections were at 2.5 mg/kg. Each animal received an intraperitoneal injection of 2.5x their body weight.

Cmp-003 solution was made by dissolving Cmp-003 (donated by Praxis Biotech) in 50% PEG400 (PanReac AppliChem, 142436.1611) and 50% sterile water. The solution was gently heated in a 40 °C water bath and vortexed every 30 s until the solution became clear. Stock Cmp-003 solution was at 0.5 mg/ml and animal injections were at 5.0 mg/kg. Solution was used immediately and made fresh daily.

Behavioral assessment of cognitive functions. For all behavioral assays the experimenter(s) were blinded to therapeutic intervention. Prior to behavioral analysis animals were inspected for gross motor impairments. Animals were inspected for whisker loss, limb immobility (included grip strength) and eye occlusions. If animals displayed any of these impairments, they were excluded. Behavioral assessment was recorded and scored using a video tracking and analysis setup (Ethovision XT 8.5, Noldus Information Technology).

Radial Arm Water Maze.

The radial arm water maze (RAWM) was used to test spatial learning and memory in rodents (26, 39). The pool is 118.5 cm in diameter with 8 arms, each 41 cm in length, and
an escape platform. The escape platform is slightly submerged below the water level, so
it is not visible to the animals. The pool was filled with water that was rendered opaque
by adding white paint (Crayola, 54–2128-053). Visual cues are placed around the room
such that they were visible to animals exploring the maze. Animals ran 6 trials a day
during learning and 3 trials during each memory probe. On both learning and memory
days there is a 10-minute inter-trial interval. Animals were trained for 2 days and then
tested on memory tests 24 hours and 7 days after training. During a trial, animals were
placed in a random arm that did not include the escape platform. Animals were allowed
1 min to locate the escape platform. On successfully finding the platform, animals
remained there for 10 s before being returned to their warmed, holding cage. On a failed
trial, animals were guided to the escape platform and then returned to their holding cage
10 s later. The escape platform location was the same, whereas the start arm varied
between trials.

Animals were injected (intraperitoneal) with either vehicle or ISRIB (2.5 mg/kg) starting
the day prior to behavior (Figure 2A) and after each of the final trials of the learning days
(day 1 and 2) for a total of three doses. No injections were given when memory was tested
on days 3 and 10. RAWM data were collected through a video tracking and analysis setup
(Ethovision XT 8.5, Noldus Information Technology). The program automatically analyzed
the number of entries into non-target arms made per trial. Every three trials were
averaged into a block to account for large variability in performance; each learning day
thus consisted of 2 blocks, whereas each memory test was one block each. Importantly,
in all animal cohorts tested (regardless of age or drug treatment) learning was measured
(Significant time effect observed in all Two-way repeated measure ANOVA analysis when
groups are analyzed independently).

Delayed Matching to Place Barnes Maze.
Beginning at day 20 animals were tested on DMP using a modified Barnes maze (26, 40).
The maze consisted of a round table 112 cm in diameter with 40 escape holes arranged
in three concentric rings consisting of 8, 16, and 16 holes at 20, 35, and 50 cm from the
center of the maze, respectively. An escape tunnel was connected to one of the outer
holes. Visual cues were placed around the room such that they were visible to animals
on the table. Bright overhead lighting and a loud tone (2 KHz, 85 db) were used as
aversive stimuli to motivate animals to locate the escape tunnel. The assay was
performed for 4 days (days 20-23). The escape tunnel location was moved for each day
and animals ran four trials on the first two days and 3 trials on the last two days. During
a trial, animals were placed onto the center of the table covered by an opaque plastic box
so they were not exposed to the environment. After they had been placed on the table for
10 s, the plastic box was removed and the tone started playing, marking the start of the
trial. Animals were given 90 s to explore the maze and locate the escape tunnel. When
the animals successfully located and entered the escape tunnel, the tone was stopped. If
the animals failed to find the escape tunnel after 90 s, they were guided to the escape
tunnel before the tone was stopped. Animals remained in the escape tunnel for 10 s
before being returned to their home cage. The maze was cleaned with ethanol between
each trial. A new escape tunnel was used for each trial. The experimenter was blind to
the treatment groups during the behavioral assay. Each trial was recorded using a video
tracking and analysis setup (Ethovision XT 8.5, Noldus Information Technology) and the program automatically analyzed the amount of time required to locate the escape tunnel. Animal improvement was calculated by Day 20 escape latency – Day 23 escape latency.

**Tissue collection.** All mice were lethally overdosed using a mixture of ketamine (10 mg/ml) and xylazine (1 mg/ml). Once animals were completely anesthetized, blood was extracted by cardiac puncture and animals were perfused with 1X phosphate buffer solution, pH 7.4 (Gibco, Big Cabin, OK, -70011-044) until the livers were clear (~1–2 min). For Western blot analysis following phosphate buffered solution (PBS), the whole brain (regions dissected discussed below) was rapidly removed and snap frozen on dry ice and stored at −80 °C until processing.

**Western Blot Analysis.** Animals received all 3 ISRIB injections and were terminated 20 h after the third injection (as described above). Frozen brain lysates or hippocampi isolates were then homogenized with a T 10 basic ULTRA-TURRAX (IKA) in ice-cold buffer lysis (Cell Signaling 9803) and protease and phosphatase inhibitors (Roche). Lysates were sonicated for 3 min and centrifuged at 13,000 rpm for 20 min at 4°C. Protein concentration in supernatants was determined using BCA Protein Assay Kit (Pierce). Equal amount of proteins was loaded on SDS-PAGE gels. Proteins were transferred onto 0.2 µm PVDF membranes (BioRad) and probed with primary antibodies diluted in Tris-buffered saline supplemented with 0.1% Tween 20 and 3% bovine serum albumin.

ATF4 (11815) (Cell Signaling) and β-actin (Sigma-Aldrich) antibodies were used as primary antibodies. HRP-conjugated secondary antibodies (Rockland) were employed to detect immune-reactive bands using enhanced chemiluminescence (ECL Western Blotting Substrate, Pierce) according to the manufacturer instructions. Quantification of protein bands was done by densitometry using ImageJ software.

ATF4 levels were normalized to β-actin expression and fold-change was calculated as the levels relative to the expression in vehicle-treated derived samples, which corresponds to 1.

**Electrophysiology.** Sagittal brain slices (250 µm) including the hippocampus were prepared from old mice (~19 mo) treated with either vehicle or ISRIB or young mice (~3 mo), treated with vehicle, 12-18 hours prior (n = 5, 7, and 5 per group respectively). Mice were anesthetized with Euthasol (0.1 ml / 25 g, Virbac, Fort Worth, TX, NDC-051311-050-01), and transcardially perfused with an ice-cold sucrose cutting solution containing (in mM): 210 sucrose, 1.25 NaH2PO4, 25 NaHCO3, 2.5 KCl, 0.5 CaCl2, 7 MgCl2, 7 dextrose, 1.3 ascorbic acid, 3 sodium pyruvate (bubbled with 95% O2 – 5% CO2, pH 7.4) (see **Supplemental Table 1 for reagent information**). Mice were then decapitated and the brain was isolated in the same sucrose solution and cut on a slicing vibratome (Leica, VT1200S, Leica Microsystems, Wetzlar, Germany). Slices were incubated in a holding solution (composed of (in mM): 125 NaCl, 2.5 KCl, 1.25 NaH2PO4, 25 NaHCO3, 2 CaCl2, 2 MgCl2, 10 dextrose, 1.3 ascorbic acid, 3 sodium pyruvate, bubbled with 95% O2 – 5% CO2, pH 7.4) at 36 °C for 30 min and then at room temperature for at least 30 min until recording.
Whole cell recordings were obtained from these slices in a submersion chamber with a
heated (32 – 34°C) artificial cerebrospinal fluid (aCSF) containing (in mM): 125 NaCl, 3
KCl, 1.25 NaH₂PO₄, 25 NaHCO₃, 2 CaCl₂, 1 MgCl₂, 10 dextrose (bubbled with 95% O₂ -
5% CO₂, pH 7.4). Patch pipettes (3–6 MΩ) were manufactured from filaments
borosilicate glass capillaries (Sutter Instruments, Novato, CA, BF100-58-10) and filled
with an intracellular solution containing (in mM): 135 KGlucose, 5 KCl, 10 HEPES, 4
NaCl, 4 MgATP, 0.3 Na₃GTP, 7 2K-phosphocreatine, and 1-2% biocytin. CA1 pyramidal
neurons were identified using infrared microscopy with a 40x water-immersion objective
(Olympus, Burlingame, CA). Recordings were made using a Multiclamp 700B (Molecular
Devices, San Jose, CA) amplifier, which was connected to the computer with a Digidata
1440A ADC (Molecular Devices, San Jose, CA), and recorded at a sampling rate of 20
kHz with pClamp software (Molecular Devices, San Jose, CA). We did not correct for the
junction potential, but access resistance and pipette capacitance were appropriately
compensated before each recording.

The passive membrane and active action potential spiking characteristics were assessed
by injection of a series of hyperpolarizing and depolarizing current steps with a duration
of 250 ms from -250 pA to 700 nA (in increments of 50 pA). The resting membrane
potential was the measured voltage of the cell 5 min after obtaining whole cell
configuration without current injection. A holding current was then applied to maintain the
neuron at -67 +/- 2 mV before/after current injections. The input resistance was
determined from the steady-state voltage reached during the -50 pA current injection. The
membrane time constant was the time required to reach 63% of the maximum change in
voltage for the -50 pA current injection. Action potential parameters including the half
width, threshold, and amplitude were quantified from the first action potential elicited.
Action potential times were detected by recording the time at which the positive slope of
the membrane potential crossed 0 mV. From the action potential times, the instantaneous
frequency for each action potential was determined (1 / inter spike interval). The
maximum firing frequency was the highest frequency of firing identified throughout all
current injections. Action potential rate as a function of current injection was examined by
plotting the first instantaneous action potential frequency versus current injection
amplitude. The F/I slope was then determined from the best linear fit of the positive values
of this plot. The action potential or spike threshold was defined as the voltage at which
the third derivative of V (d³V/dt) was maximal just prior to the action potential peak. The
action potential (AP) amplitude was calculated by measuring the voltage difference
between the peak voltage of the action potential and the spike threshold. The half-width
of the action potential was determined as the duration of the action potential at half the
amplitude. The adaptation index of each cell was the ratio of the last over the first
instantaneous firing frequency, calculated at 250 pA above the current step that first
elicited spiking. The afterhyperpolarization (AHP) was calculated as the change in voltage
from baseline (measured as the mean voltage over a 100 ms interval 600 ms after
termination of a current injection that first elicited at least 12 spikes corresponding to a
firing frequency of ~50 Hz) compared to immediately after cessation of current injection
(the minimum voltage reached in the first 175 ms immediately after cessation of current
injection). Cells were excluded from analysis if excessive synaptic input was noted during
recording of the mAHP or if the cell did not fire at least 12 spikes during current injections.
To measure the spontaneous excitatory postsynaptic currents (sEPSCs), cells were recorded in voltage clamp at a holding potential of -75 mV for 4 min, a holding potential that should have little inhibitory components given the reversal potential of chloride with these solutions. Analysis of sEPSCs was performed using a template matching algorithm in ClampFit 10.7 (Molecular Devices, San Jose, CA). The template was created using recordings from multiple pyramidal cells and included several hundred synaptic events. Access resistance (Ra) was monitored during recordings, and recordings were terminated if Ra exceeded 30 megaohms. Only stable recordings (< 50 pA baseline change) with a low baseline noise (< 8 pA root mean square) were included. The first 250 synaptic events or all the events measured in the 4 min interval from each cell were included for analysis.

Fluorescent spine imaging preparation. For fluorescent spine analysis, following PBS animals were perfused with ice-cold 4% paraformaldehyde, pH 7.5 (PFA, Sigma Aldrich, St. Louis, MO, 441244) and fixed for 4 - 24 h followed by sucrose (Fisher Science Education, Nazareth, PA, S25590A) protection (15% to 30%). Brains were embedded with 30% sucrose/ Optimal Cutting Temperature Compound (Tissue Tek, Radnor, PA, 4583) mixture on dry ice and stored at -80 °C. Brains were sectioned into 20 µm slides using a Leica cryostat (Leica Microsystems, Wetzlar, Germany) and mounted on slides (ThermoFisher Scientific, South San Francisco, CA). Slides were brought to room temperature (20 °C) prior to use. Tissues were fixed using ProLong Gold (Invitrogen, Carlsbad, CA, P36930) and a standard slide cover sealed with nail polish.

Spine density quantification. For spine density quantification, whole brains from young and old male Thy1-YFP-H transgenic line were used. 3-6 images separated by 60-140 µm in the dorsal hippocampus were imaged per animal and used for dendritic spine density analysis. 9.3 µm z-stack images were acquired on a Zeiss Laser-Scanning Confocal microscope (Zeiss LSM 780 NLO FLIM) at the HDFCCC Laboratory for Cell Analysis Shared Resource Facility. 63x magnification with a water immersion objective. All protrusions from the dendrites were manually counted as spines regardless of morphology. Two individuals (blinded to age and treatment) analyzed a total length of at least 3200 µm of dendrites from each animal using NIH FIJI analysis software (v1.52n). Individual dendritic spine was calculated as density per micron and graphed relative to old mice.

Bulk Sequencing. Frozen hippocampi were kept on dry ice and processed sample per sample until mRNA extraction. Each hippocampus was placed on wet ice and submerged with 300 µl freshly-made RNAse-free lysis buffer (20 mM Tris-HCl pH 8.0, 140 mM KCl, 5 mM MgCl2, 100 µg/ml cycloheximide, 1 mM DTT, 1% Triton X-100, 25 U/ml Turbo DNaseI (Thermo Scientific)). The tissue was then triturated for 10 strokes using a disposable RNAse-free pellet pestle, followed by 10 passages through a 2G needle. Following centrifugation for 10 mins at 4°C and 16,000 x g to remove unhomogenized debris, 50 µl of lysate was used for total RNA extraction using the DirectZol RNA miniprep kit (ZymoResearch). Lysate was mixed with 150 µl TRizol reagent (Invitrogen), mixed, and incubated at room temperature for 5 mins. After addition of 150 µl 100% ethanol, samples were further purified according to the manufacturer's instructions, including DNAsel digest, and eluted
in 30 μl RNase-free water. RNA integrity was confirmed using capillary electrophoretic analysis with BioAnalyzer RNA Pico 6000 chips (Agilent). All samples had a RIN (RNA Integrity Number) > = 9.

Next, ribosomal RNA was depleted using the RiboZero rRNA (human-mouse-rat) magnetic removal kit (Epicentre) following manufacturer’s instructions. Samples were concentrated using the RNA Clean and Concentrator 5 kit (ZymoResearch) using 1 volume of binding buffer and 1 volume 100% ethanol to ensure binding of intact, long RNA only. Samples were eluted in 8 μl RNase-free water, 1 μl was used to measure concentration using Nanodrop, and 1 μl was used to confirm efficient ribodepletion using capillary electrophoresis (BioAnalyzer RNA Pico 6000, Agilent). Sequencing libraries were synthesized using the NEBNext Ultra Directional RNA-Seq kit (NEB) according to the manufacturer’s instructions, with NEBNext Multiplex Oligos for Illumina (Index Primer Set 1, NEB) and HighPrep beads (MagBio Genomics). Depending on the concentration of each sample after adaptor ligation and purification, PCR enrichment was done for 9, 10, or 11 cycles. Final libraries were run on the BioAnalyzer (High Sensitivity DNA chips, Agilent) to ensure efficient adaptor removal and absence of overamplification. Libraries were then pooled in equimolar amounts and sequenced on a NovaSeq S4 Flow Cell.

For RNAseq data analysis, we used a combination of publicly available tools and custom scripts. Demultiplexed fastq files from different lanes were first combined, and Illumina adapter sequences were trimmed off with TrimGalore! (version_0.4.4) (www.bioinformatics.babraham.ac.uk/projects/trim_galore). Contaminating ribosomal reads were removed by mapping against a fasta file of all Mus musculus rRNA/snRNA/snRNA/MtRNA/MtRNA downloaded from the Ensembl BioMart (Ensembl 92) using STAR (v2.5.3a) (43). Quality control of raw and processed fastq files was performed on a random subset of 1 million reads per sample using FastQC version 0.11.3 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). The unmapped reads were aligned to the Mus musculus GRCm38.92 Ensembl primary assembly, again using STAR, and bam files were sorted with samtools 1.1 (44). Count generation and downstream analysis were done in R (R project v3.1.2 and v3.4.0, www.R-project.org) using a combination of the packages Rsmbreads (45), EdgeR (46) plyr, ggplot2, gplots, ggrepel, plotly, corrplot, RColorBrewer, and data.table. Low read counts were removed using the EdgeR filterByExpr function, and read counts were normalized by library size. Differential expression was determined using the EdgeR Exact Test, which allows comparison of two groups of negative binomial random variables. The false discovery rate was set to 0.05. Tests for GO and KEGG pathway enrichment in the list of differentially expressed genes were performed in R, and across multiple ontogenies and databases with DAVID 6.8 (https://david.ncifcrf.gov/) using the default settings of their Functional Annotation Clustering tool. To assess pathway enrichment in a manner that is less dependent on gene significance cutoffs, gene set enrichment analysis of the entire dataset was performed against MSigDB using the GSEA software (47, 48), ranking genes according to gene score = sign(log2FC) * (-log(adj.pvalue)), as described in Reimand et al (49). Transcription factor binding site enrichment in the first 500 bp upstream of the transcription initiation site of DEGs was done using the Cytoskape iRegulon plugin (50) with ROC threshold for AUC calculation set to 1%.
A summary of the code used for data analysis is freely available on Figshare (10.6084/m9.figshare.10007312), and sequencing data can be found in NCBI’s Sequence Read Archive under BioProject SUB6444768.

**qPCR Analysis.** Hippocampus samples, of approximately the same size per animal were process as previously described (51, 52). Relative gene expression was determined using the 2−ΔΔCt method and normalized using GAPDH. Primers used were the following:

- **CD3:** Fw 5’ TGACCTCATCGCAACTCTGCTC-3’  Rev 5’ TCAGCAGTGCTTGAACCTCAGC-3’
- **Ifit1:** Fw 5’ CTGAGATGTCACTTCACATGGAA-3’  Rev 5’ GTGCATCCCCAATGGGTTCT-3’
- **Rtp:** Fw 5’ TGGGAGCAGACATTTCAAGAAC-3’,  Rev 5’ AAGGAAAGGCCTTTTGATCCTTCAGC-3’
- **Gb10:** Fw 5’ GGAGGCTCAAGAGAAAAGTCACA-3’,  Rev 5’ AAGGAAAGGCCTTTTGATCCTTCAGC-3’
- **CCL2:** Fw 5’ GCTGACCCCAAGAAGGAATG-3’  Rev 5’ GTGCTTGAGGTGGTTGTGGA-3’
- **IL1β:** Fw 5’ TGTAATGAAAGACGGCACACC-3’,  Rev 5’ TCTTCTTTGGGTATTGCTTGG-3’
- **TNFα:** Fw 5’ TGCCTATGTCTCAGCCTCTTC-3’  Rev 5’ GAGGCCATTTGGGAACTTCT-3’
- **IL-6:** Fw 5’ TACCACTTCACAAGTCGGAGGC-3’,  Rev 5’ CTGCAAGTGCTTGAACCTCAGC-3’
- **GAPDH:** Fw 5’ AAATGGTGAAGGTCGGTG-3’  Rev 5’ TGAAGGGGTCGTTGATGG-3’

**Flow Cytometric Analysis.** To assess circulating cell populations peripheral blood was collected by cardiac puncture and transferred into an EDTA collection tube. Blood was aliquoted into flow cytometry staining tubes and stained with surface antibodies for 30-60 min at room temperature (53). Surface antibodies included anti-CD45 (FITC-conjugated; BD Biosciences), Ly-6G (PE-conjugated; BD Biosciences), CD8 (PE-Cy7-conjugated; BD Biosciences), and CD4 (APC-conjugated; BD Biosciences). Leukocyte subpopulations were identified as follows: Forward and side scatter was used to exclude debris and doublet populations. Specific T- cell populations were identified as follows: CD4 T-cell subsets were CD4+, CD45+, Ly-6G, CD8, CD11b. CD8 T-cell subsets were CD8+, CD45+, Ly-6G, CD4, CD11b. After surface antibody staining, red blood cells were lysed.
with RBC lysis (BD Biosciences). Data were collected on an LSRII (BD) and analyzed with Flowjo™ software (v10, Tree Star Inc.).

Statistics.

Figure 1B- Student’s t-test (p < 0.05). Old n = 10; Old + ISRIB n = 10.

Figure 1C- One-way ANOVA (F = 18.8, p < 0.001); with Tukey post-hoc analysis. Young n = 5, Old = 3, Old + ISRIB = 3.

Figure 2B- One-way ANOVA (F = 5.3, p < 0.05); with Tukey post-hoc analysis. Young n = 10; Old n= 25; Old + ISRIB n=21.

Figure 2C- Two-way repeated measures ANOVA revealed a significant group (p < 0.01) and time effect (p < 0.01). Old n = 18; Old + ISRIB n =16.

Figure 2D- Student t-test (p < 0.05). Old n= 18; Old + ISRIB n=16.

Figure 3B- One-way ANOVA (F = 4.461, p < 0.05); with Tukey post-hoc analysis.

Neurons: n = 10 Young (5 animals); n = 12 Old (5 animals), n = 19 Old + ISRIB (7 animals) with 1-5 neurons recorded per animal.

Figure 3E- One-way ANOVA (F = 18.57, p < 0.001) with Tukey post-hoc analysis.

Young n = 7 slides (2 animals); Old n = 12 slides (3 mice); Old + ISRIB n = 17 slides (4 mice).

Figure 4A- EdgeR Exact test, FDR 0.05.

Figure 4B- One-way ANOVA (F = 8.8; p < 0.01) with a Tukey-post analysis. Young n = 8; Old n = 7; Old + ISRIB n = 8.

Figure 4C- One-way ANOVA (F =4.2, p < 0.05) with a Tukey-post analysis. Young n = 8; Old n = 7; Old + ISRIB n = 7.

Figure 4D- One-way ANOVA (F = 12.23, p < 0.001) with a Tukey-post analysis. Young n = 8; Old n = 7; Old + ISRIB n = 8.

Figure 4E- One-way ANOVA (F = 5.2; p < 0.05) with a Tukey-post analysis. Young n = 8; Old n = 7; Old + ISRIB n = 8.

Figure 4F- Linear regression was measured by Pearson R correlation (R² = 0.27; F = 8.0, p < 0.001).

Figure 4G- Student t-test (p < 0.05). Old n = 7; Old + ISRIB n = 8.

Supplemental Table 1. List of Electrophysiology Reagents.
<table>
<thead>
<tr>
<th>REAGENTS FOR PATCHING</th>
<th>Company</th>
<th>Product#</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sucrose</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>S5016</td>
</tr>
<tr>
<td>NaH₂PO₄</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>S9638</td>
</tr>
<tr>
<td>NaHCO₃</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>S6014</td>
</tr>
<tr>
<td>KCl</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>P9333</td>
</tr>
<tr>
<td>NaCl</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>S9888</td>
</tr>
<tr>
<td>CaCl₂</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>223506</td>
</tr>
<tr>
<td>MgCl₂</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>M9272</td>
</tr>
<tr>
<td>Dextrose</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>G5767</td>
</tr>
<tr>
<td>Ascorbic acid</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>A5960</td>
</tr>
<tr>
<td>Sodium pyruvate</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>P5280</td>
</tr>
<tr>
<td>Potassium gluconate</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>P1847</td>
</tr>
<tr>
<td>HEPES</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>H3375</td>
</tr>
<tr>
<td>MgATP</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>A9187</td>
</tr>
<tr>
<td>Na₃GTP</td>
<td>Sigma-Aldrich, St. Louis, MO</td>
<td>G8877</td>
</tr>
<tr>
<td>2K-phosphocreatine</td>
<td>Millipore, Burlington, MA</td>
<td>237911</td>
</tr>
<tr>
<td>Biocytin</td>
<td>Tocris, Bristol, UK</td>
<td>3349</td>
</tr>
</tbody>
</table>
ACKNOWLEDGEMENTS

This work was supported by the generous support of the Rogers Family (to S.R. and P.W.), the UCSF Weill Innovation Award (to S.R. and P.W.), the NIH/National Institute on Aging Grant R01AG056770 (to S.R.), the NRSA post-doctoral fellowship from the NIA F32AG054126 (to K.K), the National Institute for General Medicine (NIGMS) Initiative for Maximizing Student Development (R25GM056847) and the National Science Foundation (NSF) Graduate Fellowship Program (To E.S.F), the UCSF Clinical and National Center for Advanced Translational Sciences at NIH (UCSF-CTSI Grant Number TL1 TR001871) and the NIH/NINDS (K08NS114170) (To A.N), the Programa de Apoyo a Centros con Financiamiento Basal AFB 170004 (to S.B.). P.W. is an Investigator of the Howard Hughes Medical Institute.

We thank Dr. Vikaas Sohal for providing equipment for electrophysiological recordings and advice on analysis. We thank Dr. Spyros Darmanis and Rene Sit from the Chan Zuckerberg Biohub for their assistance with analysis.

The authors would like to thank Praxis Biotech LLC, San Francisco, CA for providing samples of Cmp-003, for use in experiments described in this publication.

Microscopic imaging was obtained at the HDFCCC Laboratory for Cell Analysis Shared Resource Facility which is funded through grants from NIH (P30CA082103 and S10 ODo21818-01).

CONFLICT OF INTEREST

SB is an employee of Praxis Biotech. SB, GU and LD work at Fundacion Ciencia & Vida and receive partial funding from Praxis Biotech. P.W. is an inventor on U.S. Patent 9708247 held by the Regents of the University of California that describes ISRIB and its analogs. Rights to the invention have been licensed by UCSF to Calico. P.W. is a consultant for Praxis Biotech LLC and Black Belt TX Limited. The authors declare no other competing interests.
REFERENCES


