Microglial Remodeling of the Extracellular Matrix Promotes Synapse Plasticity

Highlights

- IL-33 is expressed by subsets of hippocampal neurons and is modulated by experience
- Microglia drive dendritic spine plasticity and memory precision via neuronal IL-33
- IL-33 gain of function mitigates some age-related decreases in spine plasticity
- Neuronal IL-33 induces microglial remodeling of the extracellular matrix

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In Brief

A form of experience-dependent neuron-microglial communication is mediated by IL-33, which promotes hippocampal dendritic spine formation, synapse plasticity, and ECM engulfment and is required for memory consolidation.
Article

Microglial Remodeling of the Extracellular Matrix Promotes Synapse Plasticity

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SUMMARY

Synapse remodeling is essential to encode experiences into neuronal circuits. Here, we define a molecular interaction between neurons and microglia that drives experience-dependent synapse remodeling in the hippocampus. We find that the cytokine interleukin-33 (IL-33) is expressed by adult hippocampal neurons in an experience-dependent manner and defines a neuronal subset primed for synaptic plasticity. Loss of neuronal IL-33 or the microglial IL-33 receptor leads to impaired spine plasticity, reduced newborn neuron integration, and diminished precision of remote fear memories. Memory precision and neuronal IL-33 are decreased in aged mice, and IL-33 gain of function mitigates age-related decreases in spine plasticity. We find that neuronal IL-33 instructs microglial engulfment of the extracellular matrix (ECM) and that its loss leads to impaired ECM engulfment and a concomitant accumulation of ECM proteins in contact with synapses. These data define a cellular mechanism through which microglia regulate experience-dependent synapse remodeling and promote memory consolidation.

INTRODUCTION

Neural circuits undergo ongoing structural changes in synaptic connectivity in response to experience. Defining the mechanisms of these remodeling processes is critical for understanding how experiences become stably encoded in the brain. Components of the innate immune system regulate synapse formation and stabilization (Bennett and Molofsky, 2019; Kaneko et al., 2008; Lee et al., 2014; Stevens et al., 2007). Consistent with this, microglia, the predominant immune cells within the brain parenchyma, have been increasingly implicated in synaptic remodeling. Microglia make ongoing and activity-dependent contact with dendritic spines (Eyo et al., 2014, 2015; Li et al., 2012; Madry et al., 2018; Tremblay et al., 2010; Wake et al., 2009). They engulf synapse components during early thalamic development and in the hippocampus (Paolicelli et al., 2011; Schafer et al., 2012; Vainchtein et al., 2018; Wang et al., 2020). However, microglia can also promote spine formation in an experience-dependent manner (Miyamoto et al., 2016; Parkhurst et al., 2013) and are required for some forms of cortical plasticity (Sipe et al., 2016).

However, the mechanisms by which these activity-dependent contacts between neurons and microglia lead to stable changes in synaptic connectivity are unknown. This is in part because molecules known to regulate microglial function are only beginning to be studied and have been challenging to examine with cell-type-specific manipulations. Our group recently identified the interleukin-1 (IL-1) family cytokine IL-33 as a novel regulator of microglial activation and function (Vainchtein et al., 2018). IL-33 is normally sequestered in the nucleus in an inactive state but, upon release into the extracellular space, is a potent activator of nuclear factor κB (NF-κB) signaling in target cells that express its obligate receptor, IL1RL1 (also known as ST2; Molofsky et al., 2015). We found that, during early postnatal development, IL-33 expressed by astrocytes in the thalamus and spinal cord drove microglial synapse engulfment and restricted excitatory synapse numbers.

Synapse remodeling continues throughout life in brain regions that change in response to experience, including the hippocampus, which is essential for learning and memory consolidation. Here we define a mechanism by which neuron-microglia signaling via IL-33 augments hippocampal plasticity in an experience-dependent manner. We found that, in the hippocampus, IL-33 was predominantly expressed by neurons, that its expression was experience dependent, and that this expression defined a molecularly distinct subset of neurons enriched for markers of...
synaptic plasticity. Conditional deletion of Il33 from neurons or its receptor from microglia led to fewer dendritic spines, decreased numbers of newborn neurons, and impaired precision of fear memory retrieval. Aging was associated with reduced neuronal IL-33 as well as deficits in memory precision, and IL-33 gain of function mitigated age-related declines in spine plasticity and neuronal activation. Finally, we found that IL-33 was necessary and sufficient for extracellular matrix (ECM) engulfment by microglia. Loss of IL-33 led to accumulation of ECM proteins, particularly around synapses and dendritic spines, and exogenous enzymatic clearance of ECM restored dendritic spine numbers in IL-33-deficient animals. These studies reveal a molecular mechanism through which microglia remodel the ECM to promote synapse plasticity and reveal a requirement for this pathway in the precise consolidation of long-term memories.

RESULTS

Experience-Dependent Regulation of Il33 Expression in Adult Hippocampal Neurons

To identify the cellular sources of IL-33 in the hippocampus, we used a nucleus-localized knockin reporter line (Il33mCherry/+; Figure 1A). Unlike in developing subcortical regions, where Il33 is primarily expressed by astrocytes (Vainchtein et al., 2018), we found that most Il33-expressing cells in the hippocampus were neurons (Figures 1A and 1B; Figure S1A). Il33 has two isoforms that produce the same protein but are regulated by alternate promoters (Talabot-Ayer et al., 2012). We found that neurons expressed the Il33b isoform, whereas astrocytes produced Il33a, suggesting that neurons may have unique regulatory mechanisms for IL-33 production (Figure S1B). Il33 was expressed by ~74% of neurons in the dentate gyrus (DG) and ~24% of neurons in the CA1 subregion (Figures 1C and 1D). Within the DG, Il33 was expressed in Prox1+ granule cells, but not Parvalbumin+ interneurons and rarely in Doublecortin+ immature granule cells (Figures S1C and S1D). We did not detect neuronal Il33 expression in the entorhinal cortex, amygdala, or thalamus, although some neurons in layer 2/3 of the prefrontal cortex also expressed Il33 (Figure S1E). These results were validated by immunostaining for IL-33 protein, which demonstrated neuronal expression that was abrogated by conditional deletion with a neuron-specific SynfCre recombinase line (Figure 1E; Figure S1F).

To determine whether Il33 expression is regulated by experience, 3-month-old Il33mCherry/+ mice were placed in an enriched environment (EE), which promotes circuit remodeling and neurogenesis (Figure 1F; Bednarek and Caroni, 2011; Bergami et al., 2015; Kempermann et al., 1997). We observed that 4 weeks of EE led to a significant increase in the proportion of Il33mCherry/+ mice and compared neurons with different levels of Il33 expression (Figure 1E). We next ranked differentially expressed genes between these subsets (Figures 2E and 2F; Table S1; Finak et al., 2015). Il33-enriched cluster 2 neurons expressed genes linked to cell adhesion, synapse assembly, and synaptic plasticity, including cadherins (Cdh10 and Cdh13; Basu et al., 2017; Duan et al., 2018). We also found enriched expression of genes related to assembly and remodeling of the ECM, including Adams9, Itga4, Col23a1, and Col25a1. We cross-referenced our data with a curated dataset of matrix-associated genes (Naba et al., 2016; Figure 2E, green dots) and identified 18 genes encoding ECM-associated proteins. Thus, Il33 expression defines a molecular subtype enriched for markers of synapse formation and ECM remodeling.

IL33 Expression Identifies a Neuronal Subset Primed for Synaptic Plasticity

Memory encoding and consolidation are circuit-specific and competitive processes (Josselyn and Tonegawa, 2020). We observed that DG neurons expressed heterogeneous levels of Il33 (Figure 1E). To determine whether this heterogeneity had functional correlates, we performed single-nucleus RNA sequencing of hippocampal neuronal nuclei (Figures 2A–2F; Figures S2A–S2E). Unbiased clustering revealed cell subsets corresponding to DG granule cells (expressing Prox1, Stxbp6, and Dock10) as well as other major neuron classes (Figure 2A; Figures S2D and S2E). The DG population further clustered into two subsets, clusters 0 and 2 (Figure 2B). To identify Il33+ subsets, we also sequenced flow-sorted Il33mCherry+ neurons and included these metadata, revealing that the majority of mCherry+ neurons overlapped with DG cluster 2 (Figures 2C and 2D; Figures S2F and 2G; STAR Methods).

We next investigated differentially expressed genes between these subsets (Figures 2E and 2F; Table S1; Finak et al., 2015). Il33-enriched cluster 2 neurons expressed genes linked to cell adhesion, synapse assembly, and synaptic plasticity, including cadherins (Cdh10 and Cdh13; Basu et al., 2017; Duan et al., 2018). We also found enriched expression of genes related to assembly and remodeling of the ECM, including Adams9, Itga4, Col23a1, and Col25a1. We cross-referenced our data with a curated dataset of matrix-associated genes (Naba et al., 2016; Figure 2E, green dots) and identified 18 genes encoding ECM-associated proteins. Thus, Il33 expression defines a molecular subtype enriched for markers of synapse formation and ECM remodeling.

We next investigated whether Il33 expression was associated with structural and functional correlates of synaptic plasticity. In the DG, granule cells recruited to encode a memory underwent a period of enhanced plasticity and spine growth that is required for proper memory consolidation (Kitamura et al., 2017; Roy et al., 2017; Ryan et al., 2015). We quantified dendritic spines in sparsely labeled mature DG granule cells in Il33mCherry/+ mice and compared neurons with different levels of Il33 expression (Figure 2G). We found that neurons with high mCherry had increased dendritic spine density compared with other subsets, as analyzed by binning (Figure 2H) or by Pearson correlation (Figure S2H). We also quantified spine head filopodia as structural markers of spine plasticity. These protrusions form prior to new spine formation or spine relocation and have been observed in the olfactory bulb and CA1 regions (Breton-Provencher et al., 2016; Orlando et al., 2012;
Figure 1. Experience-Dependent Regulation of IL-33 Expression in Adult Hippocampal Neurons

(A) Post-natal day 30 (P30) hippocampus showing Il33mCherry, Aldh1l1eGFP (astrocytes), and NeuN (neurons). Inset, DG. Scale bars, 100 μm and 20 μm (inset). ML, molecular layer; GCL, granule cell layer.

(B) Quantification of Il33-mCherry+ cells labeled with cell-type-specific markers.

(C and D) Image (C) and quantification (D) of Il33-mCherry+ neurons in the DG and CA1. Scale bar, 20 μm.

(E) IL-33 protein in DG neurons in the control (Il33fl/fl) or after neuron-specific excision of Il33 (Il33fl/fl: Synapsin1-cre). Scale bar, 10 μm.

(F) Schematic of environmental enrichment (EE) and social isolation (SI).

(G and H) Image (G) and quantification (H) of the CA1 from Il33mCherry/+ mice under standard or EE conditions. Scale bar, 20 μm.

(I) Fluorescence-activated cell sorting (FACS) gating of Il33-mCherry+ neurons from the DG and CA1.

(J and K) Quantification of Il33 expression after EE (J) or SI (K) by qPCR.

(L) Schematic of newborn neuron labeling.

(M and N) Image (M) and quantification (N) of percentage of Il33-mCherry+ newborn neurons at the indicated time points after BrdU injection (n = 3 mice/time point; ≥20 neurons/mouse). Injected at 4 weeks of age. Scale bar, 10 μm.

(O) Effect of an EE on the percentage of Il33-mCherry+ newborn neurons; BrdU injection and EE at 12 weeks, analysis at 16 weeks.

(P) Effect of SI on the percentage of Il33-mCherry+ newborn neurons; BrdU injection and SI at 4 weeks, analysis at 8 weeks.

*p < 0.05, **p < 0.01, ***p < 0.001. Statistics: two-tailed unpaired t tests. Dots represent individual mice. Data are mean ± SD. See also Figure S1.
Richards et al., 2005; Weinhard et al., 2018). We found that DG neurons with high mCherry levels had a significantly greater proportion of spines harboring spine head filopodia (Figures 2I and 2J; Figure S2I).

The expression of immediate-early genes such as Cfos and Arc identifies neuronal ensembles recruited during encoding of contextual memories (Denny et al., 2014; Liu et al., 2012). These neurons are allocated to a memory trace in a competitive...
process based on their relative excitability (Cai et al., 2016; Han et al., 2007; Park et al., 2016). To determine whether Il33-high neurons were more likely to be activated in response to learning, we placed animals in a novel environment for 1 h. We found that neuronal subsets with higher mCherry levels were more likely to express c-Fos than other subsets (Figures 2K and 2L), suggesting that these subsets may be functionally distinct. However, the total number of c-Fos+ granule cells did not differ in IL-33 conditional knockout (IL-33 cKO) mice with neuron-specific deletion of Il33 (Il33flox/flox:Syn1CreER) (Figure S2J), indicating that IL-33 does not control the size of an engram population. Rather, this could indicate that neurons with high levels of Il33 and spine remodeling represent a population more likely to engage in response to novel experiences. Altogether, our data suggest that Il33 expression is associated with a subset of neurons more likely to undergo synaptic remodeling.

**Neuron-Microglial Signaling via IL-33 Drives Experience-Dependent Spine Remodeling**

To identify cellular targets of IL-33 signaling, we quantified expression of Il11r1, which encodes its obligate co-receptor (Molofsky et al., 2015; Palmer and Gabay, 2011). Microglia were the predominant cell type expressing Il11r1 in the hippocampus, as assessed by qPCR of cell subsets isolated by flow cytometry (Figures S3A and S3B) and consistent with other regions of the CNS (Vainchtein et al., 2018). Quantitative in situ hybridization supported these results, although we also identified a small proportion of Il11r1+NeuN+ cells in the hilus (Figures 3A and 3B). Genetic deletion of Il11r1 with a myeloid-specific inducible Cre recombinase line (Il11r1flax/flax;Cx3cr1CreER/−, “IL1RL1 i-cKO”) led to a 93% ± 6% reduction in Il11r1 expression in microglia (Figures S3C and S3D). Thus, microglia are competent to respond to IL-33 signaling.

Given the strong expression of markers linked to synapse formation in Il33-enriched neurons, we next investigated whether neuronal IL-33 or microglial IL1RL1 affected dendritic spine numbers. We quantified dendritic morphology and spine density in mice with neuron-specific deletion of Il33 (Il33flox/flox;Syn1CreER, “IL-33 cKO”) using AAV9-Syn1-GFP to sparsely label hippocampal neurons (Figure 3C). We found that spine density was significantly decreased in IL-33 cKO in the DG and CA1 regions (Figure 3D; Figure S5E), although dendritic branching was largely preserved (Figure S3F). Conditional deletion of Il11r1 in microglia using IL1RL1 i-cKO mice photenocopy this decreased spine density (Figure 3E). The numbers of microglia were not altered in these mice (Figure S3G), but constitutive deletion of Il11r1 (Il11r1flox/flox;Cx3cr1Cre+/−, “IL1RL1 cKO”) led to altered microglial morphology and process coverage (Figure S3H), suggesting alterations in microglial function.

Microglial contact has been correlated with formation of new dendritic spines and induction of spine head filopodia (Miyamoto et al., 2016; Weinhard et al., 2018). We found that loss of neuronal IL-33 or microglial IL1RL1 resulted in fewer spines with spine head filopodia (Figures 3F–3H). Enrichment increased the number of spine head filopodia in control mice, whereas IL1RL1 cKO limited this increase (Figures 3I and 3J). To determine the effect of these spine alterations on synaptic function, we performed whole-cell patch-clamp recordings of DG granule cells from IL-33 cKO animals and littermate controls. We found a reduction in the frequency of miniature excitatory postsynaptic currents (mEPSCs) in IL-33 cKO mice and a trend toward increased amplitude of these mEPSCs (Figures 3K–3M; Table S2). These data are consistent with our finding of fewer dendritic spines. Taken together, these data suggest that neuron-microglia signaling via IL-33 promotes new spine formation and plasticity.

To determine whether IL-33 signaling through microglia is sufficient to drive spine formation, we next used a gain-of-function approach. To drive constitutive release of IL-33, we removed the N-terminal chromatin-binding domain containing the nuclear localization signal (NLS) and replaced it with a hemagglutinin (HA) epitope tag (Bessa et al., 2014). We then used lentiviral delivery to express this modified protein (IL-33NLSS2A-tdTomato) or a control (tdTomato) in DG neurons using a Ca2+/calmodulin-dependent protein kinase II (CaMKII) promoter (Figure 3N). We detected cytoplasmic and dendritic HA-tagged IL-33 protein, indicating that the construct was functioning as expected (Figure S3I). We found that IL-33NLSS2A was sufficient to increase dendritic spine numbers and spine head filopodia and that these effects were abrogated in IL1RL1 i-cKO mice, indicating that microglial sensing of IL-33 was required (Figures 3O–3Q). These results suggest that IL-33 acts locally to increase dendritic spines, although non-cell-autonomous effects of IL-33 release on nearby neurons are also possible. Taken together, these data indicate that IL-33 release from neurons is sufficient to promote dendritic spine formation and plasticity.

**Neuron-Microglia Signaling through IL-33 Promotes Experience-Dependent Increases in Newborn Neurons and Is Required for Remote Memory Precision**

Integration of newborn neurons in the DG requires circuit remodeling and is increased in EEs, suggesting that this process is regulated by experience (Bergami et al., 2015; Kheirbek et al., 2012; Toni et al., 2007). We tested whether neuron-microglia signaling via IL-33 affects newborn neurons by pulse-chase labeling with BrdU and quantification of label-retaining NeuN+ neurons 5 weeks later (Figure 4A; Figure S4A). Although there was no difference in the number of newborn neurons in IL-33 cKO mice at baseline, mice housed in an EE for 5 weeks had significantly fewer newborn neurons compared with controls (Figure 4B). We did not detect differences in the number of Ki67+ proliferating cells in the granule zone (Figure 4C). Additionally, we found reduced dendritic spine density in newborn neurons of Il33+/− mice (Figure S4B), indicating that synaptic integration of newborn neurons might be delayed or impaired. These results suggest that neuronal IL-33 is required for experience-dependent increases in newborn neurons.

The decreased number of newborn neurons and impaired spine plasticity prompted us to investigate whether the IL-33 pathway is involved in hippocampus-dependent memory. The DG is critical for pattern separation, a process that separates similar experiences into non-overlapping representations in the hippocampus (Miller and Sahay, 2019). Synaptic plasticity in mature granule cells plays an important role in this process (McHugh et al., 2007; Sun et al., 2020), as does newborn neuron...
Figure 3. Neuron-Microglia Signaling via IL-33 Drives Experience-Dependent Spine Remodeling

(A and B) Image (A) and quantification (B) of Il1rl1 transcript co-labeled with antibodies to cell-type-specific markers in the DG (n = 4 mice). Scale bars, 25 μm and 5 μm (inset).

(C and D) Image (C) and quantification (D) of dendritic spine density in DG granule cells sparsely labeled with AAV-Syn1-GFP in IL-33 cKO versus littermate controls (nested t test, n = 36 dendritic segments, 4 mice/genotype). Scale bars, 20 μm and 2 μm (spine inset).

(E) Spine density in IL1RL1 i-cKO versus littermate controls (nested t test, n = 27 dendritic segments, 4 mice/genotype).

(F–H) Image (F) and quantification (G and H) of percentage of spines with spine head filopodia in IL-33 cKO and IL1RL1 i-cKO versus littermate controls (arrowheads, spine head filopodia; statistics as in D and E). Scale bars, 1 μm and 0.5 μm (inset).

(I and J) Schematic (I) and quantification (J) of percentage of spines with spine head filopodia under standard or enriched conditions in IL1RL1 cKO mice versus littermate controls (2-way ANOVA, Tukey’s post hoc test, n = 18–24 dendritic segments, 3–4 mice/group).

(K–M) Representative traces (K) and quantification of mEPSC frequency (L) and amplitude (M) from IL-33 cKO mice and littermate controls (nested t test, n = 12–14 neurons, 3–4 mice/genotype).

(N) Schematic of IL-33-NLS viral gain-of-function strategy.

(O and P) Image (O) and quantification (P) of spine density after injection of control (tdTomato) or IL-33NLS virus into control or IL1RL1 i-cKO mice (one-way ANOVA, Tukey’s post hoc test, n = 29–33 dendritic segments, 3 mice/group). Scale bars, 20 μm and 1 μm (inset).

(Q) Percentage of spine head filopodia (sample sizes and statistics as in P).

*p < 0.05, **p < 0.01, ***p < 0.001. Data are mean ± SD (bar graphs) and median ± interquartile range (violin plots). Larger dots to the right of each plot indicate mean per individual mouse. See also Figure S3 and Table S2.
function (Danielson et al., 2016; Kheirbek et al., 2012; Nakashiba et al., 2012; Sahay et al., 2011). We used a contextual fear discrimination assay to assess memory and pattern separation. Mice were conditioned to a context with 3 footshocks (context A) and then exposed to the conditioned context followed by an unconditioned but partially similar context (context B). We then quantified their ability to discriminate between the two contexts 1, 14, and 28 days post-training (Figure 4D; McAvoy et al., 2016). We first tested animals with neuronal deletion of IL33, which preferentially targeted the hippocampus and prefrontal cortex, because neuronal IL-33 was not detected in other regions relevant to contextual fear memory, including the entorhinal cortex, amygdala, and nucleus reuniens (Figure S1E; Josselyn and Tonegawa, 2020; Ramanathan et al., 2018). Immediately after training, IL-33 cKO animals had intact memory recall and context discrimination (Figure 4E). However, we observed a progressive decrement in the ability of IL-33 cKO mice to discriminate contexts emerging between 14 and 28 days post-training relative to controls (Figures 4F and 4G; Figure S4C). This was largely driven by an increase in freezing behavior in the unconditioned context, suggesting generalization of the fear memory. These deficits were phenocopied in animals

Figure 4. Neuron-Microglia Signaling through IL-33 Promotes Experience-Dependent Increases in Newborn Neurons and Is Required for Remote Memory Precision

(A and B) Schematic (A) and quantification (B) of newborn neurons in IL-33 cKO mice versus controls in standard or enriched housing (n = 5 mice/group, 2-way ANOVA, Tukey’s post hoc test). Scale bars, 100 μm and 20 μm (inset). (C) Ki67+ proliferating cells in the subgranular zone (SGZ). (D) Schematic of the contextual fear discrimination assay. (E–G) Freezing in the conditioned fear context A versus unconditioned context B in control and IL-33 cKO animals 1 day (E) or 28 days (F) post-training. (G) shows the same data quantified as a memory discrimination index and includes a 14-day time point (n = 16 control and 11 IL-33 cKO mice, 2-way repeated measures [RM] ANOVA, Sidak’s post hoc tests; Figure S4C). (H–J) Freezing in context A versus context B in control and IL1RL1 cKO animals 1 day (H) or 28 days (I) post-training and corresponding discrimination indices (J) (n = 12 mice/genotype, 2-way RM ANOVA, Sidak’s post hoc tests; Figure S4D). *p < 0.05, **p < 0.01. Data are mean ± SD (B and C) and mean ± SEM (G and J). See also Figure S4.
with constitutive deletion of Il1rl1 in microglia (Figures 4H–4J; Figure S4D). We did not detect differences in anxiety-related behaviors in the open field or elevated plus maze tests in the IL-33 cKO or IL1RL1 cKO cohorts (Figures S4E–S4H). Collectively, our observations indicate that neuron-microglia communication through IL-33 is required for memory precision at remote time points, suggesting a role in memory consolidation.

**IL-33 Expression Decreases in the Aged Hippocampus and Is Associated with Decreased Memory Precision and Spine Plasticity**

Decreased synaptic plasticity and memory impairment are among the phenotypes associated with physiological aging (Wyss-Coray, 2016), which led us to investigate whether this pathway might be modulated with age. We observed a significant decline in the number of Il33+ neurons in the DG and CA1 of 18-month-old mice (Figures S5A–S5E). To determine whether aging might affect memory precision, we quantified contextual fear learning and memory precision in young (3-month-old) versus aged (18-month-old) mice, using the same fear conditioning paradigm as in Figure 4. We found that, after training, aged mice recognized the conditioned context A similarly as young mice but had impaired context discrimination and overgeneralized to the unconditioned context B (Figures S5F and S5G; Figures S5B and S5C). The motor response to shock did not differ between young and old mice, suggesting that differences in freezing were not a result of altered shock sensitivity (Figure S5D). Thus, like IL-33 cKO animals, aged mice showed reduced memory precision, although this deficit was already evident 1 day after training and persisted at a remote time point (14 days; Figure 5G). Context discrimination in this memory task tended to correlate with Il33 mRNA levels in DG granule cells (Figure S5E).

These data suggested the possibility that at least some features of cognitive aging could be affected by augmenting IL-33 signaling. Increased dendritic spine plasticity and neuronal activity are associated with improved cognition in the aged brain (Cai et al., 2016; Villeda et al., 2014) and in mouse models of Alzheimer’s disease (Roy et al., 2016). To test the effect of IL-33 on dendritic spines in the aged brain, we used the lentiviral gain-of-function approach described in Figure 3 (IL-33ΔNLS). We observed that aging was associated with a substantial decrease in the number of spines with spine head filopodia and that IL-33ΔNLS in aged mice increased the number of spine head filopodia (Figures 5H and 5I). This increase was at least partly driven by a supraphysiological increase in total spines (Figure 5J), which were not significantly decreased with age, consistent with findings by other groups (Fan et al., 2017). Aging is also associated with decreased neuronal excitability (Oh et al., 2010), and age-related deficits in cognitive flexibility are ameliorated by increasing neuronal excitability and activation (Cai et al., 2016; Villeda et al., 2014). Consistent with prior studies, we found that neuronal activation decreased with age in the DG, as measured by the percentage of labeled neurons that were c-Fos+ after exposure to a novel environment (Figures 5K–5M). IL-33ΔNLS did not affect this metric in young mice but was sufficient to increase c-Fos+ neurons in old mice to the levels observed in young animals (Figure 5M). These data suggest that aging is a physiologic context associated with decreased Il33 expression and impaired memory precision and that increasing neuronal IL-33 release in individual neurons improves some structural and functional correlates of brain aging.

**Neuronal IL-33 Drives Microglial Engulfment of the ECM**

The deficient spine numbers after loss of IL-33-IL1RL1 signaling indicated a role of IL-33 in synapse formation or maintenance and a requirement for microglia in this process. To interrogate potential mechanisms by which IL-33 affects microglia, we transcriptionally profiled hippocampal microglia after 4 h of in vivo exposure to IL-33 (Figure 6A; Figure S6A). As expected, we identified a strong activation signature after IL-33 treatment, including Il1b, Tnf, and multiple chemokines (Figure 6B). Among the top upregulated genes were the class A scavenger receptor Marco, a regulator of filopodial morphogenesis and debris clearance (van der Laan et al., 1999; Pikkarainen et al., 1999), and the adaptor protein Gas7, which is necessary for phagocytic cup formation in macrophages (Hanawa-Suetsugu et al., 2019). These findings are consistent with IL-33’s role in promoting microglial phagocytosis (Fu et al., 2016; Vainchtein et al., 2018).

We next investigated potential pathways relevant to spine formation and plasticity. Outside of the brain, IL-33 promotes tissue remodeling, which includes turnover of the ECM (Bonnans et al., 2014; Molofsky et al., 2015). Live imaging studies suggest that the brain ECM is dynamic and responsive to synaptic cues (Tønnesen et al., 2018), and multiple studies have shown that ECM proteins can restrict synaptic plasticity and spine remodeling (Bolös et al., 2018; Frischknecht et al., 2009; Oray et al., 2004; Orlando et al., 2012; Pizzorusso et al., 2006; de Vivo et al., 2013). We found that 76 genes differentially expressed in microglia after IL-33 treatment were associated with the ECM (I(naba et al., 2016; Figure 6B, green dots). Of these, we identified 9 genes classified as ECM regulators that were significantly upregulated after IL-33 treatment (Figure 6C). These included several ECM proteases (Adams4, Mmp14, Mmp25, and Cts). Interestingly, Adams4 and Mmp14 encode membrane-localized metalloproteases that predominantly cleave chondroitin sulfate proteoglycans (CSPGs), which are a major class of ECM proteins in the brain (Figure 6D; Barbolina and Stack, 2008; Kelwick et al., 2015; Levy et al., 2014).

Aggrecan is a CSPG expressed prominently in perineuronal nets throughout the brain but is also highly enriched in the diffuse ECM of the DG (Figures 6D and 6E; Figures S6B and S6C; Bolös et al., 2018; Rowlands et al., 2018). We therefore investigated whether microglia can engulf Aggrecan. Using superresolution imaging and 3D reconstruction, we identified Aggrecan immunostaining within microglia and found that it localized to CD68+ lysosomes (Figure 6F). In microglia from IL-33 cKO mice, we found decreased numbers of CD68+ lysosomes compared with controls, indicating that IL-33 promotes a phagocytic phenotype (Figures 6G and 6H). Additionally, we also found a 54% reduction in the amount of engulfed Aggrecan in microglia from IL-33 cKO mice compared with controls (Figure 6I). Conversely, IL-33 gain of function was sufficient to increase microglial process coverage, the phagocytic index, and engulfment of Aggrecan (Figures 6J and 6K; Figures 6D–S6F). In summary, we show that neuron-derived IL-33 promotes microglial phagocytosis and engulfment of the CSPG Aggrecan.
Figure 5. IL-33 Expression Decreases in the Aged Hippocampus and Is Associated with Decreased Memory Precision and Spine Plasticity

(A) Overview schematic.

(B–D) Representative flow cytometry plots (B) and quantification of the percentage of mCherry+ neurons in the DG (C) and CA1 (D) of young and old mice (t test, n = 3 mice/group).

(E) Representative image of Il33-mCherry in the DG of young and old Il33mCherry/+ mice. Scale bar, 10 μm.

(F) Contextual fear discrimination 1 day post-training in young and old mice (n = 16 young and 14 old mice, 2-way RM ANOVA, Sidak’s post hoc tests; Figure S5C).

(G) Discrimination indices 1 and 14 days post-training (statistics as in F; Figure S5C).

(H–J) Schematic and image (H) of the number of spine head filopodia (I) and total spines (J) after injections with control or IL-33 gain-of-function virus (one-way ANOVA, Tukey’s post hoc test, n = 20–27 dendritic segments, 3 animals/group). Scale bars, 2 μm and 1 μm (inset).

(K) Schematic of the neuronal activation assay after IL-33 gain of function.

(L and M) Image (L) and quantification (M) of percentage of virally labeled DG neurons that were cFos+ after 1 h in a novel environment (one-way ANOVA, Tukey’s post hoc test, 3 mice/group). Scale bar, 20 μm.

*p < 0.05, **p < 0.01, ****p < 0.0001. Data are mean ± SD (bar graphs) and median ± interquartile range (violin plots). Larger dots to the right of violin plots indicate mean per individual mouse. See also Figure S5.
Figure 6. Neuronal IL-33 Drives Microglial Engulfment of the ECM
(A) Experimental overview of microglial transcriptomics.
(B) Volcano plot of differentially expressed genes after IL-33 or PBS treatment (P_adj < 0.01). Green, ECM and ECM-associated genes (Naba et al., 2016).
(C) Genes in (B) classified as ECM regulators. Red arrows, ECM proteases.
(D) Schematic of the ECM in the DG.
(E) Aggrecan staining in the hippocampus. Scale bar, 250 μm.
(F) z stack of Iba1+ microglia in the DG co-labeled with Aggrecan and CD68+ lysosomes. Scale bar, 5 μm.

(legend continued on next page)
IL-33 Deficiency Leads to Accumulation of Perisynaptic ECM

To determine whether microglial remodeling of the ECM could affect accumulation of ECM near synapses, we analyzed Aggrecan distribution in the hippocampus. Using superresolution imaging, we found that Aggrecan in the DG molecular layer formed punctate structures that colocalized with the postsynaptic density protein Homer1. IL-33 cKO mice had a significant increase in the density of Aggrecan puncta compared with littermate controls as well as increased colocalization of Aggrecan with Homer1 (Figures 7A–7D). Brevican is another CSPG prevalent in the adult brain that can restrict synaptic plasticity (Favuzzi et al., 2017; Frischknecht et al., 2009) and is proteolytically cleaved into a smaller 53-kDa fragment by ADAMTS4 (Mayer et al., 2005; Valenzuela et al., 2014; Figure 7E). We found that IL-33 cKO mice had significantly more total Brevican and a decrease in the levels of proteolyzed Brevican compared with controls (Figure 7F–7H).

We next used IL-33 gain of function to further examine the relationship between Aggrecan and dendritic spines. We found that IL-33ΔNLS increased the number of spines devoid of Aggrecan contact and that this effect required microglial expression of the IL-33 receptor IL1RL1 (Figures 7I and 7J). In aged animals, we found a 2-fold decrease in the number of spines devoid of perisynaptic Aggrecan, consistent with studies that have found marked accumulation of ECM proteins with age (Végh et al., 2014a). IL-33ΔNLS increased the number of Aggrecan-negative spines to levels seen in young mice (Figure 7K). These data indicate that IL-33 promotes ECM remodeling and restricts ECM deposition near synapses.

Finally, we sought to determine the relationship between IL-33’s effect on ECM and dendritic spine maintenance. We injected the enzyme Chondroitinase ABC (ChABC) into the DG, which compromises ECM integrity by degrading the glycosaminoglycan side chains present on CSPGs (Figure 7L). We found that, 9 days after ChABC injection, spine density was modestly increased in controls but significantly increased in IL-33 cKO mice to the levels of ChABC-treated controls (Figures 7M and 7N). These data support the hypothesis that IL-33’s effects on spines are mediated at least in part through its effect on the ECM. Taken together, these findings directly implicate microglia as regulators of ECM clearance and spine formation.

DISCUSSION

This study defines a cytokine-mediated mechanism that acts via microglia to promote dendritic spine plasticity and precise retrieval of remote memories. We find that neurons determine microglial responses by experience-dependent regulation of IL-33, coupling neuronal state to microglial activation. Finally, we demonstrate that IL-33 drives microglial engulfment of the ECM, revealing a molecular mechanism through which microglial activation leads to increases in synaptic plasticity.

Microglia Optimize Experience-Dependent Neural Circuit Plasticity

The instructive role of IL-33 in promoting microglial functions and the fact that IL-33 expression is experience-dependent suggest two levels at which this pathway may optimize encoding. First, we show that environmental inputs, including enrichment and SI, can alter the homeostatic set point of IL33 expression in neurons. These changes occur over days to weeks and suggest that this pathway could represent a mechanism whereby experiences such as stress and deprivation affect the capacity for information storage.

Second, the heterogeneous expression of IL-33 within the neuronal population suggests that not all DG neurons are equally poised for experience-dependent remodeling. Only a fraction of granule cells recruited to an engram are reactivated during memory recall, indicating that most neurons activated during learning eventually dissociate from the memory trace (Guo et al., 2018; Kitamura et al., 2017). Thus, it is possible that engram cells expressing high levels of IL-33 could gain a competitive edge by engaging microglia that promote spine plasticity. Such a mechanism could help optimize circuit connectivity and reduce memory interference, which we found to be increased in mice deficient in IL-33-IL1RL1 signaling.

Further insights into how this mechanism might underlie diversity within engram networks would require a better understanding of the time course of IL33 expression. For example, how does new IL-33 production compare with the time course of immediate-early genes such as Cfos or Arc? We were only able to detect experience-dependent changes in neuronal IL33 over days to weeks. However, the slow turnover kinetics of the reporter used in this study limit the conclusions that can be drawn. Defining the kinetics of IL33 expression with more temporal precision and at the level of single neurons as well as defining the regulation of IL-33 protein release, which is required for its downstream effects, are important areas for future study.

Microglia Remodel the Brain ECM

How do microglial interactions with synapses promote spine formation and plasticity? It has been proposed that memories may be stored in the “pattern of holes” in the ECM (Tsien, 2013). Live imaging has shown that microglia contact dendritic spines in an activity-dependent manner. Spine head filopodium formation correlates with perisynaptic microglia contact (Weinhard et al., 2018) and can be induced by local digestion of the ECM (Orlando et al., 2012). Our data raise the possibility that these frequent microglial contacts with spines could enable microglia to locally clear or modulate the ECM around individual synapses in support of activity-dependent structural plasticity.

**Representative images (G) and quantification of CD68+ lysosomes within microglia (H) and Aggrecan protein within microglial lysosomes (I) (nested t test, n = 17–20 microglia, 4 mice/genotype). Scale bar, 5 μm and 2 μm (inset).**

**Image (J) and quantification (K) of Aggrecan in microglial lysosomes 2 weeks after infection with a control or IL-33ΔNLS virus (t test, 28 microglia, 3 mice/group). Scale bars, 25 μm and 2 μm (inset).**

*p < 0.05, **p < 0.01. Data are mean ± SEM (bar graphs) and median ± interquartile range (violin plots). Larger dots to the right of violin plots indicate mean per individual mouse. See also Figure S6 and Table S3.**
**Figure 7. IL-33 Deficiency Leads to Accumulation of Perisynaptic ECM**

(A and B) Image (A) and quantification (B) of Aggrecan punctum density in control and IL-33 cKO mice. Insets, DG ML (t test, n = 3 mice/genotype). Scale bars, 200 μm (left), 40 μm (center), 5 μm (right).

(C and D) Image (C) and quantification (D) of Aggrecan and Homer1 co-localization in the DG (circles, co-localized puncta; t test, n = 3 mice/genotype). Scale bars, 1 μm and 1 μm (inset).

(E) Brevican CSPG and proteolytic cleavage site.

(F–H) Representative western blot of Brevican (F) from hippocampal lysate in IL-33 cKO and littermate controls and quantification of uncleaved (145 kDa, G) and cleaved (53 kDa, H) forms (t test, n = 3 mice/genotype).

(I and J) Aggrecan co-localization with dendritic spines in the DG injected with control (tdTomato) or IL-33 gain-of-function (IL-33 ΔNLS) virus in control or IL1RL1 i-cKO mice (one-way ANOVA, Tukey’s post hoc test, n = 29–33 dendritic segments, 3 mice/group). Scale bar, 1 μm.
Notably, the ECM is not only relevant to synapses. For example, oligodendrocytes produce ECM, which is required for proper assembly of nodes of Ranvier (Susuki et al., 2013). New myelin formation also promotes long-term memory precision (Pan et al., 2020; Steadman et al., 2020). This raises the question of whether microglia remodel myelin or other components of the extracellular space.

**IL-33 Function in Physiology, Aging, and Pathology**

Dysregulation of innate immune homeostatic mechanisms and of microglia are strongly implicated in the cognitive dysfunction that occurs in brain aging and disease (Bennett and Molofsky, 2019; Kelley et al., 2018). The progressive decline in neuronal IL-33 expression with age coincides with defects in memory precision and accumulation of perisynaptic ECM (Végh et al., 2014a). Interestingly, clearance of ECM and exogenous recombinant IL-33 are beneficial in mouse models of Alzheimer’s disease (Fu et al., 2016; Végh et al., 2014b). Thus, precisely defining the molecular mechanisms, such as IL-33, that regulate microglial synaptic function in the healthy brain may lead to new tools and approaches for cognitive repair.

**STAR+METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.cell.2020.05.050.

**ACKNOWLEDGMENTS**

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**AUTHOR CONTRIBUTIONS**


**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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**REFERENCES**


(K) Aggrecan co-localization with spines in young and old mice injected with a control or IL-33-NLS virus (one-way ANOVA, Tukey’s post hoc test, n = 19–27 dendritic segments, 3 mice/group).
(L and M) Schematic and image of Wisteria floribunda agglutinin (WFA) staining after ChABC digestion in the DG (L), (M) shows WFA localization around DG granule cells. Dashed circles, DG. Scale bars, 500 µm (L) and 20 µm (M).
(N) Dendritic spine density in IL-33 cKO or control mice injected with ChABC or vehicle (2-way RM ANOVA, Sidak’s post hoc test, n = 3 mice/genotype; dots, means per mouse; lines connect vehicle- and ChABC-injected hemispheres; violin plots show distribution of dendritic segments per group, 23–30 segments). *p < 0.05, **p < 0.01, ***p < 0.001. Data are mean ± SD (bar graphs) and median ± interquartile range (violin plots). Larger dots to the right of violin plots indicate means per mouse.


## STAR★METHODS

### KEY RESOURCES TABLE

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## Software and Algorithms

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## RESOURCE AVAILABILITY

### Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Anna Molofsky (anna.molofsky@ucsf.edu).
Materials Availability
Plasmids generated in this study will be deposited to Addgene.

Data and Code Availability
Single nuclei RNA sequencing and microglia bulk RNA sequencing data were deposited in NCBI’s Gene Expression Omnibus (Barrett et al., 2013) under accession number GSE150714.

EXPERIMENTAL MODELS AND SUBJECT DETAILS

Mice
All mouse strains were maintained in the University of California San Francisco specific pathogen–free animal facility, and all animal protocols were approved by and in accordance with the guidelines established by the Institutional Animal Care and Use Committee and Laboratory Animal Resource Center. Littermate controls were used for all experiments when feasible, and all mice were back-crossed > 10 generations on a C57BL/6 background unless otherwise indicated. The mouse strains used are described in the Key Resources Table and as referenced in the text. The following strains were used: IL-33 reporter line Il33H2B-mCherry (Vainchtein et al., 2018); Astrocyte reporter line Aldh1l1eGFP (Gong et al., 2003; Molofsky et al., 2014); Synapsin-Cre transgenic mice for neuron-specific excision of IL-33, cre expression was maintained in the female line for breeding (Knoferle et al., 2014; Rempe et al., 2006; Zhu et al., 2001); Il1rl1flox/flox and Il33fl/fl (Chen et al., 2015); Cx3cr1creER (used in studies referring to ‘IL1RL1 i-cKO’; (Yona et al., 2013) Cx3crr cre (used for studies referring to ‘IL1RL1 cKO’; (Yona et al., 2013)). All experiments, including behavioral analyses incorporated animals of both sexes in approximately equal numbers. All experiments of n > 5 were analyzed for sex-specific trends, and none were evident unless specifically noted.

Environmental enrichment consisted of housing 2-3-month old mice in large ventilated rat cages (210 sq. in, Lab Products) with standard nesting, rodent foraging toys, and one running wheel per cage (https://larc.ucsf.edu/rodent-enrichment). Social isolation conditions consisted of singly housing 4-week old mice immediately after weaning in standard mouse cages devoid of enrichment toys.

Viral Constructs
For sparse labeling of hippocampal neurons, adeno-associated virus (AAV9-hSyn1-eGFP) was packaged and supplied by the Addgene/UPENN vector core (Addgene viral prep # 105539-AAV9) at titers of ~1x10¹³ vg/mL. For IL-33 gain of function experiments, lentiviral vectors (LV-CaMKII-tdTomato and LV-CaMKII-IL33(DNLS)-2A-tdTomato) were designed in house and packaged by the UCSF virus core at titers of ~1x10⁷ vg/mL. For labeling adultborn neurons, MMLV retrovirus vector (CAG-GFP) was supplied by Addgene (Plasmid #16664) and packaged into retrovirus by the UCSF virus core at titers of ~1x10⁶ vg/mL. LV-Synapsin-GFP lentivirus was purchased from SignaGen Laboratories.

METHOD DETAILS

Tamoxifen and BrdU injections
Tamoxifen (Sigma, T5648) was diluted in Corn Oil (Sigma-Aldrich) at 37°C overnight and administered by oral gavage at a concentration of 200 mg/kg four times every other day beginning at 4 weeks of age. For all BrdU labeling experiments, BrdU (Sigma-Aldrich) was diluted in a 10 mg/mL stock solution in sterile 1X PBS and injected intraperitoneally at a concentration of 100 mg/kg daily for 3 days.

Stereotaxic injections
All brain injections were performed with a Kopf stereotaxic apparatus (David Kopf, Tujunga, CA) with a microdispensing pump (World Precision Instruments) holding a beveled glass needle with ~50 µm outer diameter. For perinatal injections to sparsely label hippocampal neurons, P2-3 mice were anesthetized by hypothermia, headfixed with a custom clay mold, and 0.5 µL of AAV9-hSyn1-eGFP was injected at a rate of 15 nL/sec into the right ventricle using in house coordinates (1.5 mm AP from lambda, 0.8 mm ML, −1.6 mm DV). A dilution of 1:250 in sterile saline was used for experiments measuring dendritic spine density and a 1:750 dilution was used for experiments requiring single neuron morphology. Mice were allowed to recover on a heat pad before being returned to their cage.

For injections into adult hippocampus, mice were anesthetized with 1.5% isoﬂurane at an oxygen flow rate of 1L/min, headfixed with a stereotaxic frame, and treated with ophthalmic eye ointment. Fur was shaved and the incision site was sterilized with 70% ethanol and Betadine prior to surgical procedures. A hole was drilled in the skull and 1.5 µL of LV-CaMKI1-ttdTomato or LV-CaM-KI1-IL33(ANLS)-2A-ttdTomato was injected at a 1:25 dilution in sterile saline (−2.1 mm AP, 1.375 mm ML, x −1.7 mm DV) at a rate of 3 nL/sec. For injections with Chondroitinase ABC (Sigma), 40 U/mL were injected with LV-Synapsin-GFP (SignaGen Laboratories) at a dilution of 1:150. Body temperature was maintained throughout surgery using a heating pad. Post-surgery, Buprenorphine (Henry Schein Animal Health) was administered as needed by intraperitoneal injection at a concentration of 0.1 mg/kg.
Immunohistochemistry and Confocal Microscopy

For all immunohistochemistry experiments, mice were perfused transcardially with ~10 mL of ice-cold 1X PBS followed by ~10 mL of 4% (weight/volume) paraformaldehyde diluted in PBS. Brains were post-fixed in 4% PFA for a minimum of 4 hours and then transferred to a 30% sucrose solution for a minimum of 2 days. Brains were then flash frozen and sliced in 40 um thick coronal sections on a HM 440E freezing microtome (GMI Instruments).

For immunohistochemistry, brain sections were incubated in a blocking solution consisting of 10% normal goat or donkey serum (Thermo Fisher) and 0.4% Triton (Sigma-Aldrich) diluted in 1X PBS. Primary antibodies were diluted in 3% normal goat or donkey serum in 0.4% Triton and tissue was incubated on a shaker overnight at 4°C. Secondary antibodies were diluted in 3% normal goat or donkey serum and tissue was incubated on a shaker for 2 hours at room temperature with Hoechst 33342 (Thermo Fisher) added at 1:5000 during the last 5 minutes of incubation. Brain sections were mounted on coverslips with ProLong Gold without DAPI (Thermo Fisher) for high resolution imaging and ProLong Gold (Thermo Fisher) for all other experiments.

For IL-33 staining, brain sections were heated in a 1X Sodium Citrate solution (Fisher Scientific) at 95°C for 10 minutes and then washed 3x with PBS prior to blocking. For BrdU staining, brain sections were stained with non-BrdU primary antibodies and secondary antibodies as described above and then incubated with 4% PFA for 10 minutes at room temperature. Brain sections were incubated in 2N HCl (Sigma-Aldrich) for 30 minutes at 37°C and then quenched with Boric Acid buffer (0.31 g/50 mL in H2O, pH 8.5) for 10 minutes at room temperature before being carefully washed four times with 1X PBS. Tissue was incubated in a blocking solution of 10% normal goat serum and 0.4% Triton on a shaker for 1 hour at room temperature and then incubated with an anti-BrdU antibody (Abcam) diluted in 3% normal goat serum and 0.4% Triton on a shaker overnight at 4°C. Secondary staining and mounting were performed as described above.

Histology slides were imaged on an LSM 880 confocal microscope with Airyscan (Zeiss) on Superresolution mode using a 63x objective for high resolution imaging and an LSM 700 confocal microscope (Zeiss) using 20x and 63x objectives for all other imaging experiments.

Fluorescent In Situ Hybridization (FISH)

FISH experiments were performed using the RNAscope Multiplex Fluorescent Reagent Kit v2 assay (ACD Bio) as described by the manufacturer for fixed-frozen tissue, except tissue was not baked for 30 minutes at 60°C prior to tissue dehydration. Brains were embedded in OCT following 30% sucrose treatment and frozen at ~80°C for a minimum of 1 day and then sliced in 15 um thick coronal sections on a CryoStar NX70 Cryostat (Thermo Fisher) before being mounted on coverslips for downstream RNAscope processing. Mouse Il1rl1 RNAscope Probe (ACD Bio, 440661) and TSA Plus Cyanine 3 reagent (Perkin Elmer) were used to detect Il1rl1 transcript. For immunohistochemical labeling with antibodies following the RNAscope assay, tissues were incubated with blocking and antibody solutions as described above immediately after developing the HRP-TSA Plus signal and carefully washing four times.

Flow cytometry

For all flow cytometry experiments, animals were perfused transcardially with ice-cold 1X PBS and hippocampi were isolated under a dissecting microscope where meninges and excess myelin were removed. Antibodies are outlined in the Key Resource Table. Gating for neuronal and antibody solutions as described above immediately after developing the HRP-TSA Plus signal and carefully washing four times.

For sorting neuronal nuclei, cells were isolated using a protocol modified from Jaegar et al., 2018. Hippocampi were isolated under a dissecting microscope and mechanically dissociated using a 2 mL glass tissue homogenizer (Kontes Glassware) on ice using five strokes of the loose pestle and ten strokes of the tight pestle in nuclei isolation media (250 mM sucrose, 25 mM KCl, 5 mM MgCl₂, 10 mM Tris pH 8.0, and nuclease free H₂O). Dissociated nuclei were pelleted in an eppendorf tube at 1000 g for 5 minutes at 4°C and then resuspended in nuclei storage buffer (166.5 mM sucrose, 5 mM MgCl₂, 10 mM Tris pH 8.0, and nuclease free H₂O) using a P1000 tip. Nuclei were again pelleted at 1000 g for 5 minutes at 4°C and resuspended in blocking buffer consisting of 1% (weight/volume) RNase-free BSA and 1X PBS. Cells were filtered and then pelleted at 300 g for 5 minutes at 4°C before being resuspended in 22% Percoll (GE Healthcare) and centrifuged at 900 g for 20 minutes with acceleration set to 4 and deceleration set to 1 in order to remove cellular debris. Pelleted microglia were then washed with isolation medium and incubated in block solution consisting of anti-mouse CD16/32 antibody (BioLegend). Cells were sorted on a BD FACSAria III and gated on forward/side scatter, live cells by Hoechst, CD11b-PE (eBioScience), and CD45-FITC (BioLegend) to isolate microglia.

For sorting multiple CNS cell-types in parallel for comparative qPCR, cells were isolated and fixed as described (Srinivasan et al., 2016). Hippocampi were roughly minced with a sterile razor blade, incubated in 1X Accutase (Millipore) at 4°C for 1 hour, pelleted at Cell 182, 388–403.e1–e7, July 23, 2020 e4.
2000 g for 1 minute, and resuspended in Hibernate A Low Fluorescence medium (BrainBits). Cells were then lightly pelleted using a benchtop microcentrifuge and supernatant filtered through a 70 μm cell strainer into a FACS tube. 1 mL of Hibernate A media was then added to the cell pellet and slowly triturated with a P1000 tip. Cells were again lightly pelleted and supernatant was filtered into FACS tubes. This process was repeated 2 additional times. To remove excess myelin, cells were then resuspended in 22% Percoll diluted with Hibernate A and centrifuged at 900 g for 20 minutes with acceleration set to 4 and deceleration set to 1. To fix and permeabilize cells, cell pellets were resuspended in 500 uL of Hibernate A and 500 uL of ice cold 100% ethanol was slowly added. Cells were incubated on ice for 15 minutes, washed twice with Hibernate A, and then incubated in block solution consisting of anti-mouse CD16/32 antibody (Biolegend). Cells were sorted on a BD FACS Aria III and gated on forward/side scatter, intact cells by Hoechst, CD11b (eBioscience) to identify microglia, GFAP (Millipore) to identify astrocytes, Olig2 (Thermo Fisher) to identify oligolineage cells, and NeuN (Millipore) to isolate neurons.

Single nuclei RNA-sequencing

Neuronal nuclei were isolated from hippocampus, stained with Alexa Fluor 488 mouse anti-NeuN 1:1500 (Millipore), and sorted on an Aria III (BD) flow cytometer. To account for “dropouts,” we also isolated NeuN+ neuronal nuclei from Il33mCherry/+ mice and sorted the top 30% of mCherry+ neurons by intensity and sequenced them as metadata. FACS purified neuronal nuclei were sequenced using the Chromium single cell gene expression platform (10x Genomics, version 2). Approximately 15,000 nuclei from each sample were loaded into each well of Chromium Chip A according to the manufacturer instructions and combined into droplets with barcoded beads using the Chromium controller. Libraries were prepared by the Gladstone Institutes Genomics Core following the instructions in the Chromium Single Cell 3’ Reagent Kits version 2 user guide. The samples were sequenced to an average depth of 40,000–60,000 reads on an Illumina HiSeq 4000 sequencer.

Single nuclei data analysis

Sequenced samples were processed using the Cell Ranger 2.1 pipeline and aligned to the GRCm38 (mm10) mouse reference genome. In order to count the numerous intronic reads aligned from our nuclear dataset, a custom pre-mRNA GTF file was made using cellranger by converting the “transcript” annotation in the reference genome to “exon,” enabling CellRanger to count pre-mRNAs. Clustering and differential expression analysis were conducted using Seurat version 2.3.4 (Butler et al., 2018; Satija et al., 2015). Cells with fewer than 500 detected genes/cell or over 3000 genes/cell, comprising the lower and upper 5% of cells respectively, or more than 1% mitochondrial DNA were excluded during quality control steps before identification of variable genes in the dataset, cell centering, and scaling. Additionally, 3000 genes were eliminated that were expressed by fewer than 5 cells in the dataset.

After log normalization and scaling the data with a scale factor of 10,000, approximately 1500 of the most variable genes were identified and used for calculating nearest neighbors, PCA, and t-SNE plots. The most variable genes were identified using a LogVMR function from the Seurat package with parameters of expression mean between 0.125 and 5, and dispersion > 0.5. 40 PCs were calculated using these genes and the first 15 principle components were used for subsequent calculations. Clustering and t-SNE plots were computed using Seurat (Butler et al., 2018; Satija et al., 2015). A resolution of 0.5 was used to generate clusters, and the clusters were named based on expression of canonical marker genes for each region of the hippocampus. For differential expression analysis of the dentate gyrus, cells from the two clusters identified in the dentate gyrus (0, 2) were extracted from the dataset, re-normalized, and scaled. The top 3000 most variable genes were calculated using the LogVMR method in Seurat; only genes with mean expression between 0.1 and 5 and dispersion over 0.1 were included in subsequent calculations. The cells were then clustered as described for the full dataset. Cells were identified as “female” or “male” based on their expression of the gene Xist; any cells expressing one or more counts of Xist were labeled female, while all others were labeled male. Female cells from sample 3 were known to express IL-33 based on IL-33mCherry fluorescence during sorting prior to sequencing. Differentially expressed genes were identified using the “FindMarkers” function in Seurat, comparing the cluster enriched in sorted IL-33 expressing cells with the other dentate gyrus cell cluster. Only genes expressed in 25% of the cells in a given cluster that passed a log fold change threshold of 0.25 were included in the differentially expressed gene list. GO analysis was conducted using the Metascape webpage (https://www.metascape.org).

Bulk RNA sequencing

Microglia were isolated from hippocampus, stained with CD45-FITC 1:100 (BioLegend) and CD11b-APC 1:100 (BioLegend), and sorted on an Aria III (BD) flow cytometer into RLT plus buffer (QIAGEN). RNA was extracted and purified using an RNaseq Kit (QIAGEN). RNA quality was measured using the Agilent RNA 6000 Pico kit on an Agilent Bioanalyzer. All samples had RNA Integrity Number (RIN) > 7. For each sample, 10 ng of RNA was used as input for cDNA amplification and library construction using the Quant-Seq 3’ mRNA-Seq Library Prep Kit FWD for Illumina (Lexogen) following manufacturer’s instructions. Library quality and quantity was assessed using the Agilent High Sensitivity DNA kit on an Agilent Bioanalyzer and Quant-iT dsDNA Assay Kit on a Qubit Fluorometer (Thermo Fisher Scientific). Single-end (65-bp reads) sequencing was performed using an Illumina HiSeq 4000 yielding 30–45 million reads per sample.
Bulk RNA sequencing analysis
Quality of reads was evaluated using FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc), and all samples passed quality control, and reads were aligned to mm10 (GRCm38; retrieved from Ensembl, version November 2019) using STAR (version 2.5.4b) (Dobin et al., 2013) with ‘–outFilterMultimapNmax 1’ to only keep reads that map one time to the reference genome. Mapped reads were counted using HTSeq (version 0.9.0) (Anders et al., 2015) and DESeq2 package (version 1.24.0) (Love et al., 2014) was used to normalize the raw counts and perform differential gene expression analysis (using the apeglm method (Zhu et al., 2019) for effect size shrinkage). Volcano plot was made using genes from the comparison IL-33 versus PBS with adjusted p value < 0.01 with the EnhancedVolcano package (version 1.2.0).

Behavioral Assays
Conditioned fear was elicited by administering three mild footshocks (0.75 mA) following a five-minute exposure to an array of contextual cues (conditioning chamber, chamber lights, white noise, scent of anise extract). Retrieval of the fear memory was assessed by a three-minute re-exposure of the animal to the conditioning context in the absence of shock and freezing (the cessation of all movement outside of respiration) was interpreted as expression of fear memory. Video recordings were acquired and scored automatically in FreezeFrame (Actimetrics). Mice were habituated to transport and holding in a separate room for at least one hour prior to all conditioning or retrieval sessions and subjected to three days of handling prior to all behavioral tests. For assessment of fear discrimination, freezing was measured in a context similar to the conditioning context two hours later, but with the following variations: the chamber fan was turned off, the scent of anise extract was swapped for lemon extract, and a plastic divider was inserted to make the chamber walls circular and opaque. Freezing in the similar context was tested two hours following retrieval testing in the original conditioning context, and animals were rested in a holding room between sessions.

Anxiety-like behaviors were assessed via ten-minute exposures to the open field test (OFT) and five minute exposures to the elevated plus maze (EPM). Video recordings were acquired in Ethovision (Noldus) and time spent in the periphery of the OFT and in the closed arms of the EPM were scored through automated object detection and interpreted as anxiety-like behavior. Freezing and anxiety-like behaviors were both scored via an automated, unbiased process and all behavioral testing and analysis was performed blinded to genotype.

Slice Preparation and Patch-Clamp Electrophysiology
After euthanizing mice with 4% isoflurane and removing brains, 300 mm-thick transverse slices were prepared in ice-cold sucrose cutting solution (234 mM sucrose, 2.5 mM KCl, 1.25 mM NaH2PO4, 10 mM MgSO4, 0.5 mM CaCl2, 26 mM NaHCO3, and 10 mM glucose, equilibrated with 95% O2 and 5% CO2, pH 7.4) using a Leica VT1200 microtome (Leica Microsystems). Hippocampal sections were incubated for an hour at 32-34°C and then at room temperate in artificial cerebrospinal fluid (aCSF; 126 mM NaCl, 2.5 mM KCl, 1.25 mM NaH2PO4, 1 mM MgCl2, 2 mM CaCl2, 26 mM NaHCO3, and 10 mM glucose, equilibrated with 95% O2 and 5% CO2, pH 7.4).

Recording electrodes were made from borosilicate glass with a resistance 4–5.5 MΩ when filled with intracellular solution (115 mM potassium gluconate, 11 mM KCl, 1 mM MgCl2, 1 mM CaCl2,10 mM HEPES, 11 mM EGTA, and 2 mM K2ATP, pH adjusted to 7.35 with KOH; 286 mM Osm). Series resistance was monitored in all recordings, and the recordings were excluded from analysis if the series resistance was > 33 MΩ or varied by more than 10%. Recordings were obtained using a MultiClamp 700B microelectrode amplifier (Molecular Devices, Sunnyvale, CA), digitized using Digidata 1550B (Molecular Devices), and acquired at 20 kHz using pClamp 10 software (Molecular Devices). Recordings were performed in voltage-clamp mode at a holding potential of ~70 mV and obtained from visually identified mature granule cell neurons in the mid and outer granule cell layer of the dentate gyrus, which were confirmed by morphology and intrinsic membrane properties. In the presence of aCSF supplemented with 0.5 μm tetrodotoxin (TTX) and 50 μm picrotoxin, miniature excitatory post-synaptic currents (mEPSCs) were isolated and recorded for 5 minutes. mEPSCs were analyzed using ClampFit (Molecular Devices) and MiniAnalysis (Synaptosoft) software. Experimenter was blinded to genotype.

qPCR
To extract RNA from cells isolated by FACS, freshly sorted cells were pelleted at 500 g for 10 minutes at 4°C and then resuspended in RLT Plus buffer (QIAGEN). Cells were vortexed and frozen for at least one day at –80°C before being thawed on ice and processed for RNA using an RNeasy Mini Kit (QIAGEN). Purified mRNA was converted to cDNA with the High Capacity cDNA Reverse Transcription kit (Life Technologies) and amplified using either the Fast SYBR Green Master Mix (Thermo Fisher) or TaqMan Gene Expression Master Mix (Thermo Fisher) and a 7900HT Fast Real-Time PCR System (Applied Biosystems).

Western blotting
Hippocampi were dissected from freshly perfused mouse brains and excess meninges and myelin removed under a dissecting microscope. Tissues were then flash frozen on dry ice and sonicated for 20 s in lysis buffer (50 mM tris-HCl, 1 mM EDTA, 150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate and 0.1% SDS). The sample was centrifuged for 10 minutes at 15,000 rpm at 4°C and the pellet was discarded. Samples were run on a denaturing gel and transferred to PVDF membrane, blocked with 5% milk in TBST for 1 hour at room temperature, incubated with Brevican antibody (BioLegend, MMS-5127) at 1:200 overnight at 4°C and secondary at room temperature for one hour, and developed with ECL plus.
**BrdU quantification**

BrdU+ NeuN+ cells in the granule cell layer in the DG were counted in every sixth coronal hemibrain section and the total from six sections were multiplied by six to estimate the number of adultborn neurons.

**Dendritic Spine and Spine Head Filopodia quantification**

Images were acquired with an LSM 880 Confocal Microscope with Airyscan (Zeiss) on Superresolution mode using a 63x objective (NA 1.4) with 5x optical zoom (0.03 um pixel size). Z stacks were acquired at a step-size set at half the optical section and laser power and gain were consistent across all experiments. Airyscan processing was performed in Zen software (Zeiss) at a setting of 6 (“optimal” setting). Images were taken in the molecular layer of the DG. Dendritic spine density was quantified using ImageJ by measuring a distance of at least 15 μm along a dendritic branch and then counting dendritic spines. The experimenter was blinded to genotype for all quantifications. For experiments comparing IL-33 cKO, IL1RL1 cKO, and IL1RL1 i-cKO mice to controls, AAV9-Syn1-GFP virus was injected into P2-3 pups intracerebroventricularly at a dilution of 1:250 in sterile saline and mice were analyzed at 2-4 months of age.

For experiments stratifying dendritic spines and spine head filopodia by Il33-mCherry expression, the soma of sparsely labeled granule cells were imaged using a 20x objective with 16-bit depth and the position of each neuron was recorded. Dendritic spines for each neuron were imaged at 63x as described above. Mean fluorescent intensity for mCherry was then quantified in the nucleus using ImageJ and mCherry negative neurons were identified manually. For mCherry+ neurons, Il33-mCherry low and high neurons were designated as the bottom and top half of neurons by mCherry intensity, respectively. For these experiments, AAV9-Syn1-GFP virus was injected into P2-3 pups intracerebroventricularly at a dilution of 1:750 in sterile saline and mice were analyzed at 2 months of age.

**Scholl analysis**

Z stacks were acquired from 60 μm thick coronal sections containing hippocampus with an LSM 700 Confocal microscope (Zeiss) using a 20x objective. Scholl analysis was performed in ImageJ using the simple neurite tracer plugin (https://imagej.net/SNT).

**Aggrecan and Homer1 puncta quantification**

Images were acquired with an LSM 880 Confocal Microscope with Airyscan on Superresolution mode using the same specifications as described above for dendritic spines. Aggrecan puncta were quantified using the Puncta Analyzer plugin (Ippolito and Eroglu, 2010) in ImageJ and thresholding was consistent for every image. For colocalization of Aggrecan and Homer1 puncta, images were analyzed with the Puncta Analyzer plugin with a minimum pixel specification of 4. Three images in the molecular layer were averaged per mouse for analysis.

**Aggrecan engulfment**

Z stacks encompassing entire microglia were collected on an LSM 880 Confocal Microscope with Airyscan (Zeiss) on Superresolution mode using a 63x objective (NA 1.4) with 5x optical zoom (0.03 um pixel size); laser power and gain were consistent across all experiments. Images were analyzed using Imaris software (Bitplane) by creating a 3D surface rendering of individual microglia, which were thresholded to ensure microglia processes were accurately reconstructed and maintained consistent thereafter. The microglia rendering was used to mask the CD68 channel within that microglia, which was then 3D rendered, and the CD68 volume was subsequently used to mask the Aggrecan channel. Aggrecan engulfment per microglia was then calculated as the total volume of Aggrecan within the masked CD68 volume.

**Microglia coverage**

Images of Iba1 immunostaining were acquired on an LSM 700 Confocal microscope (Zeiss) using a 20x objective with 16-bit depth. Microglia coverage was quantified in ImageJ by first thresholding the Iba1 channel and then measuring the area of Iba1+ pixels and normalizing that value by the total area per field of view in the DG molecular layer. Thresholding was held consistent for each image.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Graphpad Prism 8.3.0 was used for most statistical analyses. Statistical tests are as described in text and figure legends. Violin plots were used for data with n ≥ 20 to better visualize the distribution of individual data points. RNA-sequencing data was analyzed in R as described in the methods section above.
Figure S1. IL-33 Expression in the Adult Brain at Baseline and after Enrichment, Related to Figure 1
(A) Il33 expression (Il33-mCherry) in hippocampal astrocytes (Aldh1l1-GFP) and oligolineage cells (Olig2). Scale bar: 10 μm.
(B) Representative gating and expression of Il33 transcript isoforms from qPCR of flow sorted hippocampal astrocytes and dentate granule cells (t test, n = 4 mice).
(C) DG neurons with immunolabeling for Il33-mCherry, the granule cell marker Prox1, and interneuron marker parvalbumin (PV). Scale bar: 20 μm.

(legend continued on next page)
(D) DG neurons with immunolabeling for Il33-mCherry, NeuN, and the progenitor/immature neuron marker Doublecortin (Dcx). Scale bar: 10 μm.

(E) Expression of Il33-mCherry with the astrocyte reporter Aldh1l1-eGFP and NeuN across brain regions in adult mice. Scale bar: 50 μm. 3V: third ventricle.

(F) Immunostaining for IL-33 protein and neuronal labeling with AAV9-Syn1-GFP. Scale bar: 5 μm.

(G and H) Schematic and time course of the percentage of Il33-mCherry+ neurons in the CA1 region after indicated time periods of environmental enrichment quantified by flow cytometry (2-way ANOVA, Tukey’s post hoc test, n = 3-4 mice/group).

(I) Proportion of Il33-mCherry+ neurons in DG and CA1 after 4 weeks of enrichment versus standard housing quantified by flow cytometry (t test; n = 4 mice).

(J) Proportion of Il33-mCherry+ neurons after 4 weeks of social isolation versus standard housing quantified by flow cytometry (t test; n = 4 mice).

(K and L) Percentage of NeuN+ neurons that are Il33-mCherry+ after 4 weeks of enrichment versus standard housing in the CA2 region, as demarcated by RGS14 staining, and CA3 (t test; n = 3 mice/condition). Scale bar: 200 μm.

*p < 0.05, **p < 0.01, ***p < 0.001. Data = mean ± SD.
**Figure S2. Quality Control and Validation of snRNA-Seq, Related to Figure 2**

(A and B) Quality control metrics for single nuclei RNA sequencing. (A) Number of genes recovered per cell. (B) Mitochondrial RNA content per cell. Lines show thresholds used for cell selection prior to all other analyses.

(C) t-SNE plot showing clustering of the independent wild-type replicates (coral and turquoise).

(D and E) Violin and t-SNE plots of markers used to assign subtype identities (Figure 2B) showing genes for interneurons (Gad2), CA3 pyramidal neurons (Socs2), DG granule cells (Prox1), CA1 pyramidal neurons (Satb2), and CA2/CA region pyramidal neurons (Sv2b).

(F) Percentage of cells per cluster originating from WT (blue) and IL33-mCherry (red) samples.

(G) Gating strategy to isolate neuronal nuclei and IL33-mCherry positive subsets.

(H and I) Correlation between mCherry levels and dendritic spine or spine head filopodia numbers (Pearson correlation, n = 20 neurons, > 110 spines per neuron, 3 mice). Different colors correspond to neurons from individual mice.

(J) Representative image and quantification of the number of cFos+ cells in the DG of IL-33 cKO and control mice (t test, n = 4 mice/genotype). Scale bar: 200 μm.
Figure S3. Additional Data Pertaining to IL1RL1 Expression and IL-33 Loss- and Gain-of-Function Experiments, Related to Figure 3
(A) Gating strategy for isolation of hippocampal microglia (CD11b), neurons (NeuN), astrocytes (GFAP), and oligodendrocytes (Olig2) by FACS and expression of cell-type specific markers by qPCR (n = 4 mice).
(B) Expression of Il1rl1 transcript in neurons and glia by qPCR of flow sorted hippocampal cell types (n = 3 mice). Scale bar: 50 µm.
(C) Schematic and representative image of Il1rl1 excision specificity in the hippocampus using the Cx3cr1creER allele fate mapped with the inducible Rosa26-tdTomato reporter (Ai14) and immunostaining with Iba1 and NeuN (n = 3 mice). Scale bar: 50 µm.
(D) Sensitivity of Il1rl1 excision by qPCR of flow sorted microglia using IL1RL1 i-cKO mice (Cr3cr1CreER/+:Il1rl1flox/flox; one-way ANOVA; Tukey’s post hoc test, n = 3 mice/group).
(E) Representative image of CA1 pyramidal neuron and quantification of dendritic spine density after loss of neuronal IL-33 (t test, n = 30-34 dendritic segments, 3 mice/genotype). Scale bar: 20 µm.
(F) Scholl analysis of dendritic branching in DG granule cells after loss of neuronal IL-33 (t test, Holm-Sidak multiple comparisons; n = 12-16 neurons, 3 mice). Scale bar: 20 µm.
(G) Number of Iba1+ microglia in the DG of IL-33 cKO and control mice (t test, n = 3-4 mice). Scale bar: 200 µm.
(H) Representative images of Iba1+ microglia from control and IL1RL1 cKO mice housed under standard or enriched conditions (4 weeks). Quantification of the percentage of Iba1+ pixels (microglia coverage) in the molecular layer of the DG (one-way ANOVA, Tukey’s post hoc test, n = 3-4 mice/group). Scale bar: 20 µm.
(I) Representative image of HA epitope tag immunostaining after viral transduction with control (tdTomato) or IL-33 gain-of-function (IL-33ΔNLS) constructs. Scale bar: 20 µm.
*p < 0.05, **p < 0.01, ***p < 0.001. Data = mean ± standard deviation (bar graphs) and median ± interquartile range (violin plots).
Figure S4. Additional Data Pertaining to Neurogenesis and Behavior, Related to Figure 4

(A) Representative image of BrdU labeled neurons 35 days post-injection and quantification of the percentage of BrdU+ label-retaining cells that are NeuN+ in IL-33 cKO and control mice (t test, n = 5 mice/genotype). Scale bar: 20 μm.

(B) Dendritic spine density in 4-week old newborn neurons labeled with MMLV-GFP retrovirus, which selectively targets proliferating cells, in IL-33 KO and control animals (t test, n = 34-53 dendritic segments, 3 mice). Scale bar: 20 μm, 2 μm (inset).

(C) Statistics for contextual fear discrimination assay in IL-33 cKO mice versus controls (related to Figures 4E–4G)

(D) Statistics for contextual fear discrimination assay in IL1RL1 cKO mice versus controls (related to Figures 4H–4J)

(E and F) Open field and Elevated plus maze testing of IL-33 cKO mice versus controls (t test, n = 16 control mice, 11 IL-33 cKO mice).

(G and H) Open field and Elevated plus maze testing of IL1RL1 cKO mice (t test, n = 12 mice/genotype).

*p < 0.05. Data = mean ± SD.
Figure S5. Additional Data Pertaining to Behavior and IL-33 Expression during Aging, Related to Figure 5

(A) Representative gating strategy to isolate neuronal nuclei from Prox1+Ctip2+ dentate granule cells and Prox1–Ctip2+ pyramidal neurons with fluorescence minus one (FMO) negative staining controls.

(B and C) Quantification of freezing in context A versus context B fourteen days post-training in young and old mice (n = 16 young mice and 14 old mice, 2-way RM ANOVA, Sidak’s post hoc tests). Statistics for contextual fear discrimination assay in young versus old mice (also related to Figures 5F and 5G).

(D) Quantification of distance traveled in young and old mice during the first 2 s footshock (t test, n = 16 young and 14 old mice. Mean ± SD).

(E) Correlation between day 14 discrimination index and expression of Il33 mRNA in flow sorted dentate granule cells in individual old mice (Pearson correlation, n = 11 mice).

*p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001.
Figure S6. Additional Data on Microglial Transcriptomics and Aggrecan Engulfment, Related to Figure 6
(A) Representative gating strategy to isolate live microglia (CD11b+CD45+).
(B) Representative hemisection from adult (P90) mouse showing Aggrecan immunostaining. Inset = hippocampus. Scale bar: 500 μm, 250 μm (inset).
(C) Aggrecan colocalization with WFA+ perineuronal nets around Parvalbumin+ interneurons in CA1. Scale bar: 50 μm.
(D and E) Representative image and quantification of microglial process coverage index calculated as the percentage of Iba1+ pixels in a field of view normalized to the percentage of tdTomato+ pixels after viral transduction with control or IL-33ΔNLS constructs (t test, n = 3 mice). Scale bar: 20 μm.
(F) Quantification of number of CD68+ lysosomes in microglia after viral transduction with control or IL-33ΔNLS constructs (t test, 28 microglia, 3 mice/group).
*p < 0.05. Data = mean ± standard deviation (bar graph) and median ± interquartile range (violin plot).