Interdependence of neural network dysfunction and microglial alterations in Alzheimer’s disease-related models

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Epileptiform activity and microglial mRNA changes in aged hAPP mice

Antiepileptic drug treatment and tau ablation reduce both abnormalities

TREM2 reduction exacerbates epileptiform activity after excitotoxin challenge

Neural network and microglial dysfunctions could form a vicious circle in AD
Interdependence of neural network dysfunction and microglial alterations in Alzheimer’s disease-related models

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SUMMARY
Nonconvulsive epileptiform activity and microglial alterations have been detected in people with Alzheimer’s disease (AD) and related mouse models. However, the relationship between these abnormalities remains to be elucidated. We suppressed epileptiform activity by treatment with the antiepileptic drug levetiracetam or by genetic ablation of tau and found that these interventions reversed or prevented aberrant microglial gene expression in brain tissues of aged human amyloid precursor protein transgenic mice, which simulate several key aspects of AD. The most robustly modulated genes included multiple factors previously implicated in AD pathogenesis, including TREM2, the hypofunction of which increases disease risk. Genetic reduction of TREM2 exacerbated epileptiform activity after mice were injected with kainate. We conclude that AD-related epileptiform activity markedly changes the molecular profile of microglia, inducing both maladaptive and adaptive alterations in their activities. Increased expression of TREM2 seems to support microglial activities that counteract this type of network dysfunction.

INTRODUCTION
Alzheimer’s disease (AD), the most prevalent and costly neurodegenerative disorder (Wimo et al., 2017; El-Hayek et al., 2019; GBD 2016 Dementia Collaborators, 2019; Alzheimer’s Association, 2020), is associated with diverse forms of neural network dysfunction, including non-convulsive epileptiform activity (Vossel et al., 2013, 2016; Palop and Mucke, 2016; Horváth et al., 2017; Lam et al., 2017, 2020; Lam and Noebels, 2020). However, the causes and consequences of AD-associated network dysfunctions remain to be fully elucidated. Evidence obtained in AD-related mouse models suggests that the human amyloid precursor protein (hAPP) or some of its metabolites and the microtubule-associated protein tau (MAPT) causally contribute to these dysfunctions (Palop and Mucke, 2016; Harris et al., 2020; Johnson et al., 2020; Chang et al., 2021; Hector and Brouillette, 2021). Indications that hypersynchronous network activity may promote disease progression include the observations that cognitive decline is faster in AD patients with detectable epileptiform activity than those without (Vossel et al., 2016), that treatment with the antiepileptic drug levetiracetam (LEV) reverses functional magnetic resonance imaging abnormalities and reduces some cognitive deficits in people with amnestic mild cognitive impairment (Bakker et al., 2012, 2015), and that treatment with LEV, or its analog brivaracetam, suppresses epileptiform activity and reverses both synaptic deficits and cognitive impairments in different lines of hAPP transgenic mice (Sanchez et al., 2012; Shi et al., 2013; Nygaard et al., 2015; Fu et al., 2019). hAPP mice also simulate multiple other features of AD, including pathologically elevated levels of amyloid-β (Aβ) peptides, amyloid plaques, dystrophic neurites, astrogliosis, and microgliosis (Palop and Mucke, 2016; Johnson et al., 2020). The molecular and cellular mechanisms that link neural network dysfunctions to other AD-relevant alterations and pathogenic processes have yet to be defined.

Independent lines of evidence have implicated myeloid cells and microglial dysfunction in the pathogenesis of AD, but the roles of microglia in AD have been studied primarily in relation to genetic risk factors and pathological hallmarks such as amyloid plaques and neurofibrillary tangles (Meyer-Luehmann and Prinz, 2015; Efthymiou and Goate, 2017; Sarlis and Heneka, 2017; Hammond et al., 2019; Leng and Edison, 2021). For example, variants of the triggering receptor expressed on myeloid cells 2 (TREM2) markedly increased expression of TREM2 seems to support microglial activities that counteract this type of network dysfunction.
increase AD risk and have been shown to modulate both amyloid and tangle pathology in mouse models (Guerreiro et al., 2013; Jonsson et al., 2013; Gratuze et al., 2018; Ulland and Colonna, 2018). Other studies demonstrated that experimental enhancement of specific brain oscillations changes the morphology of microglia, expression of cytokines, and amyloid load in brain tissues of hAPP mice (Iaccarino et al., 2016; Martorell et al., 2019; Garza et al., 2020). However, it is unclear whether aberrant neural network activities that spontaneously arise in AD or related models alter microglial gene expression and whether preventing or reversing such neuronal dysfunctions can prevent or reverse altered immune responses in AD-relevant contexts. In the current study, we investigated the impact of non-convulsive epileptiform activity on the expression of microglial and other inflammation-related gene products in hAPP mice from line J20 (Mucke et al., 2000; Palop and Mucke, 2016; Johnson et al., 2020; Gulbranson et al., 2021) as well as the impact of TREM2 hypofunction on excitotoxin-induced network hypersynchrony.

**RESULTS**

**LEV treatment suppresses neural network dysfunction and reverses inflammation-related gene expression changes in aged hAPP-J20 mice**

It is widely thought that alterations in the expression of inflammation-related genes in brains of patients with AD and of related animal models are primarily caused by structural pathological alterations, particularly amyloid plaques and neurofibrillary tangles (Meyer-Luehmann and Prinz, 2015; Sarlus and Heneka, 2017; Gratuze et al., 2018; Leng and Edison, 2021). To determine whether functional alterations in neural network activities also contribute to gene expression changes by innate immune cells in AD-relevant contexts, we treated 22–26-month-old, plaque-laden (Figure 1A) male heterozygous hAPP-J20 mice and age/male sex-matched, nontransgenic, wildtype (WT) controls from the same line with placebo or LEV for four months using chow formulations (Figure 1B). Intracranial EEG recordings at 2 and 4 months after treatment initiation revealed abnormally increased epileptiform activity in placebo-treated hAPP-J20 mice, as compared with placebo-treated WT controls (Figures 1C–1D). LEV treatment suppressed the epileptiform activity in hAPP-J20 mice, although it did not eliminate it completely at the dosage used here (Figures 1C–1D). Consistent with results obtained after treatment with LEV or the closely related drug brivaracetam in younger hAPP-J20 and APP/PS1 mice (Sanchez et al., 2012; Nygaard et al., 2015; Fu et al., 2019), LEV treatment reduced behavioral abnormalities in a replicate cohort of aged hAPP-J20 mice, including hyperactivity in the open field (Figure 1E) and spatial learning and memory deficits in the active place avoidance paradigm (Figures 1F–1H).
To identify altered immune responses, we used a NanoString Neuroinflammation panel to compare hippocampal mRNA levels in the four groups of mice that had undergone EEG recordings while they were treated with LEV or placebo. Compared with placebo-treated WT mice, more inflammation-related transcripts were increased than decreased in placebo-treated hAPP-J20 mice (Figure 1I; Table S1). LEV treatment changed the expression of multiple inflammation-related genes in hAPP-J20 mice (Figure 1J) and the majority of these changes reversed alterations that differentiated hAPP-J20 mice from WT controls (Figures 1K–1L).

To look for potential interactions among transcripts whose levels were altered by both hAPP expression and LEV treatment, we used STRING analysis (Szklarczyk et al., 2019). This approach provides information on direct (physical) and indirect (functional) associations derived from genomic context, co-expression data, text-mining, biochemical/genetic experiments, and previously curated pathway and protein-complex databases. Of the 25 gene products whose elevated levels in hAPP-J20 mice were reduced by LEV treatment, 20 showed interactions (Figure 1M). The majority of genes in this cluster are predominantly expressed in microglia and macrophages, and several are implicated in AD or related conditions. For example, the complement factor C1Q is found at abnormally high levels in brains of AD patients and hAPP mice; it is concentrated near amyloid plaques and colocalizes with synapses, possibly tagging them for elimination (Hong et al., 2016; Sarlus and Heneka, 2017; Schartz and Tenner, 2020). C1Q levels are also elevated in epileptic conditions not associated with AD pathology (Schartz et al., 2018; Andoh et al., 2019). Expression of the pattern recognition receptor and immune response mediator CLEC7A is increased in microglia/macrophages in brains of hAPP mice (Keren-Shaul et al., 2017). Variants of TREM2 strongly increase AD risk (Guerreiro et al., 2013; Jonsson et al., 2013; Efthymiou and Goate, 2017). Levels of the TYRO protein tyrosine kinase-binding protein (TYROBP), an adaptor protein for TREM2, are increased in AD brains, and variants of TYROBP increase the risk for early-onset AD (Zhang et al., 2013; Pottier et al., 2016).

Quantitative RT-PCR (qRT-PCR) analysis revealed similar hAPP- and LEV-dependent modulations of these AD-relevant transcripts in the cortex of the mice whose hippocampal RNA we had analyzed with the NanoString Neuroinflammation panel (Figures 1N–1Q), demonstrating that the effects of our experimental manipulations are not restricted to one brain region. This set of data also illustrates how effectively LEV treatment counteracts the increased expression of these transcripts in hAPP-J20 mice, even though it does not affect their levels in WT controls (Figures 1N–1Q).

Some of the transcripts most strongly altered by both APP expression and LEV treatment were expressed predominantly by microglia/macrophages associated with amyloid plaques, as demonstrated by colabeling of coronal hemibrain sections from 26 to 30-month-old hAPP-J20 and WT mice with mRNA probes, antibodies against the cell-specific marker IBA1, and the amyloid stain Thioflavin S (ThioS) (Figure 2). In hAPP-J20 mice, C1qa, Trem2, and Tyrobp mRNA signals colocalized with IBA1-positive cells and the highest expression levels were seen around ThioS-positive plaques (Figures 2A–2C). None of the respective probes yielded detectable signals in WT mice (data not shown). Thus, at least some of the genes that were altered by hAPP and LEV treatment are expressed in microglia or infiltrating macrophages.

We also measured C1Q immunoreactivity in brain sections from hAPP-J20 and WT mice after LEV and placebo treatment to confirm that the changes we found in C1qa, C1qb, and C1qc transcript levels result in commensurate changes in C1Q protein levels. Compared to age-matched WT mice, C1Q immunoreactivity was increased within the hippocampus of hAPP-J20 mice, particularly in the molecular layer of the dentate gyrus and the stratum lacunosum-moleculare of the CA1–CA3 subfields (Figure S1). LEV treatment normalized hippocampal C1Q immunoreactivity levels in hAPP-J20 mice (Figure S1).

The most parsimonious interpretation of these results is that the microglia/macrophage-related gene expression changes that LEV treatment reduced in hAPP-J20 mice are caused, at least in good part, by epileptiform activity. The following experiments were designed to further test this hypothesis and to exclude alternative possibilities.

**LEV treatment does not modulate amyloid plaques and plaque-associated microgliosis in aged hAPP-J20 mice or LPS-induced gene expression changes in other models**

Previous studies demonstrated that treating 3-month-old hAPP-J20 mice with LEV or 13-month-old APP/PS1 mice with brivaracetam for 3–4 weeks did not alter Aβ levels or amyloid loads in their brain tissues.
To assess whether the more prolonged LEV treatment used here affected amyloid plaques and plaque-associated microgliosis in aged mice, we analyzed coronal hemibrain sections from 26–30-month-old hAPP-J20 mice after they had been treated for 4 months with LEV or placebo (Figure 1B). ThioS staining showed that both groups of mice had robust amyloid deposition in the hippocampus (Figure 3A). Quantitations of amyloid plaque numbers, plaque-associated ThioS signal intensities, plaque areas, and plaque circularities revealed no differences between LEV- and placebo-treated hAPP-J20 mice (Figures 3B–3E). In a similar vein, quantitative analyses of brain sections co-labeled with ThioS and an antibody against IBA1 revealed no differences between these groups of mice in regard to the number and morphology of microglia/macrophages associated with their plaques (Figures 3F–3I).

While the above findings make it unlikely that LEV treatment affects the expression of inflammation-related genes by changing plaques or the physical association between plaques and microglia, another possibility...
Figure 3. LEV does not alter amyloid plaque loads or plaque-associated microgliosis in hAPP-J20 mice
(A–I) Coronal brain sections from 26–30-month-old hAPP-J20 mice that had been treated with LEV or placebo (Figure 1) were stained with ThioS (magenta) to label plaques and with TO-PRO3 to label nuclei (cyan) (A–E) or with ThioS (magenta) and an IBA1 antibody to label microglia/macrophages (blue) (F–I). (A) Representative images of the hippocampus. Scale bars: 500 μm, 50 μm (inset). (B) Number of plaques per mm² of hippocampal area. (C) Fluorescence intensity of ThioS-positive plaques. Mean levels of fluorescence intensity in placebo-treated hAPP-J20 mice were defined as 1.0. (D) Area covered by ThioS-positive plaques. (E) Plaque circularity. A perfect circle would equal 1.0. (F) Representative images of microglia/macrophages. Scale bar: 50 μm. (G) Number of IBA1-positive cell bodies within 25 μm of a ThioS-positive plaque. (H) Fraction of IBA1-positive pixels that were also positive for ThioS. (I) Soma size (area) of IBA1-positive cells that were located > 25 μm (distal, Dist.) or < 25 μm (adjacent, Adj.) of a ThioS-positive plaque. n = 7 mice per group (3 sections were analyzed per mouse) for (B–E and G–I). For (G–I), three plaques were analyzed in each section. For (I), three microglia distal to a plaque and three microglia adjacent to a plaque were analyzed. Unpaired, two-tailed Student’s t-test of the data in (B–E and G–I) revealed no significant differences between groups. Two-way ANOVA of the data in (I) revealed an effect of distance to plaques (p < 0.001), but no significant effect of LEV treatment (p = 0.41) and no interaction between these variables (p = 0.52). *p < 0.05, **p < 0.01 vs. leftmost bar or as indicated by brackets by two-way ANOVA and Holm-Sidak test. Dots in (B–E and G–I) represent individual mice. Bars are means ± s.e.m.
to consider is that LEV might have direct effects on the expression of such genes. To explore whether LEV has immune-modulatory effects that are independent of its anti-epileptic activities, we activated the immune system in vivo by injecting 5–6-month-old male WT mice intraperitoneally (IP) with lipopolysaccharide (LPS), a manipulation that causes inflammatory responses throughout the body (Sousa et al., 2018), and in vitro by adding LPS to the medium of microglial cultures. In mice, LPS did not increase the frequency of epileptiform spikes (Figures S2A–S2B), but it robustly altered the expression of inflammation-related genes in the cortex (Figures S2C–S2H). Some of the gene expression changes pointed in opposite directions from those observed in placebo-treated hAPP-J20 mice. For example, C1qa and Trem2 transcript levels were decreased in LPS-treated WT mice (Figures S2C–S2D), consistent with previous findings (Sousa et al., 2018), but increased in placebo-treated hAPP-J20 mice (Figures 1N and 1P). Such differential responses are not surprising, as it is well established that microglia and macrophages react to distinct pathological challenges with differential changes in their molecular profiles (Flaris et al., 1993; Sousa et al., 2018). Most importantly, pre-treating mice with LEV did not change the LPS-induced alterations or affect the expression of inflammation-related genes in placebo-treated mice (Figures S2C–S2H).

LPS treatment also elicited robust gene expression changes in cultures of primary microglia from WT mice (Figures S2I–S2N). Treating the cultures with LEV over a range of doses did not modulate the LPS-induced changes or affect microglial gene expression under baseline (vehicle control) conditions (Figures S2I–S2N). Thus, LEV does not appear to directly affect microglial gene expression in vivo or in vitro.

Early LEV treatment or tau ablation suppresses neural network dysfunction and inflammation-related gene expression changes in hAPP-J20 mice

To assess the relationship among inflammation-related gene expression changes, epileptiform activity, and much earlier stages of amyloid deposition, we tested whether suppressing epileptiform activity could reverse inflammation-related gene expression changes in hAPP-J20 mice that were just beginning to form plaques in the hippocampus and had not yet formed plaques in the cortex (Mucke et al., 2000; Sanchez et al., 2012; Fu et al., 2019; Whitesell et al., 2019) and data not shown). Starting at 6 months of age, hAPP-J20 mice and WT controls were treated with LEV or placebo for 1 or 3 months. Hippocampal and cortical levels of C1qa, Trem2, and Tyrobp mRNAs were determined by qRT-PCR at the end of the treatments. Compared with placebo-treated WT controls, increases in the expression of these transcripts were mostly seen in the hippocampus (Figure 4). At both ages, LEV treatment of either length reduced the increased expression of C1qa, Trem2, and Tyrobp mRNAs in hAPP-J20 mice but did not affect their levels in WT controls (Figure 4).

Next, we prevented the development of epileptiform activity in hAPP-J20 mice through the genetic ablation of endogenous mouse tau, a strategy that effectively counteracts network hypersynchrony in mouse models with epilepsy of diverse causes, including in hAPP-J20 mice (Roberson et al., 2007, 2011; Chang et al., 2021). EEG recordings in 22–26-month-old male and female Mapt+/+ and Mapt–/– mice that did or did not express hAPP confirmed that hAPP-J20/Mapt+/+ mice had an increased frequency of epileptiform spikes and that tau ablation prevented this abnormality in hAPP-J20/Mapt–/– mice (Figure 5A). Analysis of hippocampal mRNA from these mice by NanoString Neuroinflammation panel revealed that more inflammation-related transcripts were increased than decreased in hAPP-J20/Mapt–/– mice, as compared to WT controls, and that transcripts expressed predominantly in microglia and macrophages featured prominently among those that differed between these genotypes (Figure 5B; Table S2).

Tau ablation prevented 43 gene expression changes in hAPP-J20 mice, including increased expression of 28 and decreased expression of 15 transcripts (Figures 5C–5D; Table S2). In the absence of hAPP, tau ablation changed the expression of only one of the genes we investigated, Mapt (Table S2). To look for potential interactions among transcripts whose levels were altered by both hAPP and tau, we again used STRING analysis (Szklarczyk et al., 2019). Of the 28 gene products whose elevated levels in hAPP-J20 mice were reduced by tau ablation (Figures 5E), 25 showed interactions (Figure 5F). The majority of genes in this cluster are predominantly expressed in microglia and macrophages, and several have been implicated in AD or related conditions, similar to the cluster of genes whose aberrant expression in hAPP-J20 mice was reversed by LEV treatment (Figure 1M).

qRT-PCR analysis confirmed hAPP- and tau-dependent modulations of AD-relevant microglial transcripts in the cortex (Figures 5G–5J), again demonstrating that the effects of our experimental manipulations are...
not restricted to the hippocampus in aged mice. Tau ablation effectively prevented the increased expression of \( \text{C1qa} \), \( \text{Clec7a} \), \( \text{Trem2} \), and \( \text{Tyrobp} \) mRNAs in hAPP-J20 mice but did not affect the levels of these transcripts in mice without hAPP expression (Figures 5G–5J). We previously demonstrated that tau ablation does not alter A\( \beta \) levels or the extent of amyloid pathology in brains of hAPP-J20 mice (Roberson et al., 2007). Thus, reduction of endogenous tau prevents epileptiform activity as well as aberrant gene expression by microglia/macrophages in the presence of pathologically elevated levels of hAPP/A\( \beta \).

Figure 4. Effects of LEV treatment on microglial gene expression changes in young hAPP-J20 mice

(A–L) Male hAPP-J20 mice and WT controls were treated for 4 or 12 weeks with LEV (+; 2 g/kg of chow) or placebo (−) starting at 6 months of age. At the end of treatments, hippocampal and cortical mRNA was analyzed by qRT-PCR. (A–F) Hippocampal (A–C) and cortical (D–F) levels of the indicated transcripts after 4 weeks of LEV or placebo treatment. (G–L) Hippocampal (G–I) and cortical (J–L) levels of the indicated transcripts after 12 weeks of LEV or placebo treatment. Mean levels of the respective transcripts in placebo-treated mice without hAPP expression were defined as 1.0. \( n = 4–5 \) mice per group. *\( p < 0.05 \), **\( p < 0.01 \), ***\( p < 0.001 \), ****\( p < 0.0001 \) vs. leftmost bar or as indicated by brackets (two-way ANOVA and Holm-Sidak test). Two-way ANOVA revealed a significant effect of genotype on all transcripts but one (F, \( p = 0.18 \); E,K,L, \( p < 0.05 \); A,D, \( p < 0.01 \); C, \( p < 0.001 \); B,G,H,I,J, \( p < 0.0001 \)), a significant effect of LEV treatment on some transcripts (B,E,H,I, \( p < 0.05 \); G, \( p < 0.01 \)) but not others (A,C,D,F,J–L, \( p > 0.05 \)), and a significant interaction between these variables for some transcripts (I, \( p < 0.05 \); D, \( p < 0.01 \)) but not others (A–C,E–H,J–L, \( p > 0.05 \)). Dots represent individual mice and bars \( \pm \) s.e.m.
Notably, several of the inflammation-related gene expression changes that were prevented by tau ablation in aged hAPP-J20 mice were also reversed by LEV treatment (Figures 5K–5M; Table S3). Furthermore, multiple gene expression changes correlated with the frequency of epileptiform spikes in untreated aged male and female hAPP-J20 mice and in placebo-treated aged male hAPP-J20 mice. Expression changes of 13 genes correlated with the frequency of epileptiform spikes in at least two of these groups (Figure 6A), and Tyrobp expression levels correlated with the frequency of epileptiform spikes in all three groups (Figures 6B–6D). Taken together, these results support the hypothesis that the effects of tau ablation on inflammation-related gene expression in hAPP-J20 mice resulted from the prevention of epileptiform activity.

Hypofunction of TREM2 enhances epileptiform activity

Although inflammatory changes, and microglial activities in particular, are often suspected of promoting the development of disease, the immune system clearly also fulfills many protective functions (Wyss-Coray and Mucke, 2002; Meyer-Luehmann and Prinz, 2015; Sarlus and Heneka, 2017; Schartz and Tenner, 2020; Leng and Edison, 2021). TREM2 is especially interesting in this regard, as its hypofunction seems to be a strong risk factor for AD (Guerreiro et al., 2013; Jonsson et al., 2013; Efthymiou and Goate, 2017; Ulland and Colonna, 2018) and its expression was robustly increased in hAPP-J20 mice and normalized by both LEV treatment (Figures 1I, J, 1L, 1P, 4B, and 4H) and tau ablation (Figures 5B, 5C, 5E, and 5I). We therefore wondered whether TREM2 has a role in microglial responses to neural network dysfunction.

To explore this possibility and simulate the hypofunction of TREM2 that is thought to increase AD risk in heterozygous carriers of certain TREM2 variants (Guerreiro et al., 2013; Jonsson et al., 2013; Efthymiou and Goate, 2017; Cheng-Hathaway et al., 2018; Sudom et al., 2018; Ulland and Colonna, 2018), we analyzed heterozygous Trem2 knockout (Trem2+/−) mice, whose TREM2 levels are reduced by 50%. To directly address whether hypofunction of TREM2 modulates epileptiform activity, we challenged Trem2+/− and WT mice with a subthreshold dose (10 mg/kg of body weight) of the proepileptic drug kainate, an approach that avoids confounding interactions between TREM2 and hAPP metabolites. Although we detected no difference in epileptiform activity between Trem2+/− mice and WT (Trem2+/+) controls at baseline, Trem2+/− mice had more epileptiform activity than WT mice after the kainate challenge (Figures 7A–7C). Longitudinal analysis of EEG recordings revealed that after the kainate injection, peak spike frequencies were much higher and reached faster in Trem2+/− than Trem2+/+ mice (Figures 7D–7F). The excitotoxin-induced epileptiform activity also took much longer to subside in Trem2+/− than Trem2+/+ mice, resulting
in an increased overall burden of epileptiform spikes in Trem2+/− mice (Figures 7G–7I). Taken together, these findings suggest that the increased expression of TREM2 in hAPP-J20 mice represents an adaptive response, an important purpose of which may be to curtail hypersynchronous neural network activity.

DISCUSSION

The results from our investigation of hAPP-J20 mice suggest that the aberrant expression of inflammation-related genes in brains with pathologically elevated levels of hAPP/Aβ and extensive plaque loads are driven and mediated, at least in part, by epileptiform activity, a type of neural network dysfunction that also occurs in patients with AD (Vossel et al., 2013, 2016; Palop and Mucke, 2016; Horváth et al., 2017; Lam et al., 2017, 2020; Lam and Noebels, 2020). This conclusion is supported by our findings that these gene expression changes correlated with the extent of epileptiform activity and could be reduced by two distinct experimental interventions (LEV treatment and tau ablation) that suppress epileptiform activity in aged, plaque-laden hAPP-J20 mice. Pearson correlation coefficients (r) and p values are shown in the graphs. Dots represent individual mice.

Although LEV treatment and tau ablation markedly reduced inflammation-related gene expression changes in aged hAPP-J20 mice, it remains possible that these gene expression changes are not caused solely by epileptiform activity, but also depend on the presence of amyloid plaques, which could prime
Figure 7. Genetic reduction of Trem2 exacerbates epileptiform activity after low-dose kainate challenge

(A–I) EEG recordings were obtained from 6-month-old male Trem2+/− mice and WT (Trem2+/+) controls before and after a single IP injection of kainate (10 mg/kg body weight). (A) Schematic of experiment. (B) Sample EEG traces during the first hour after kainate injection. (C) Spike frequency 24 h before and after kainate injection. (D) Number of spikes per min during the first hour after kainate injection. (E) Peak spike frequency after kainate injection. (F) Time to reach peak spike frequency after kainate injection. (G) Change in spike frequency over 35 days after kainate injection. (H) Cumulative spikes estimated by calculating the area under the curve in (G). (I) Kaplan–Meier curves indicating the percentage of mice with abnormally increased epileptiform activity relative to their own baseline at different days after the kainate injection. *p < 0.05, **p < 0.01, ***p < 0.001 vs similarly treated WT mice or as indicated by the bracket, based on two-way ANOVA and Holm-Sidak test (C) or unpaired, two-tailed Student’s t test (E, F, and H). Values in (C–H) are means ± s.e.m.
Taken together, these findings indicate that TREM2 critically contributes to microglial functions that counteract hypersynchronous network activity and hippocampal amyloid plaques, but only sparse Aβ deposits in the cortex (Johnson et al., 2020). Furthermore, our in situ hybridization experiments showed that these gene expression changes in hAPP-J20 mice were most prominent in microglia/macrophages associated with plaques.

Since the main target of LEV is the synaptic vesicle glycoprotein 2A (Lösch et al., 2016) and we did not observe any direct effects of LEV on microglia in primary cultures lacking neurons, we find it most plausible to hypothesize that the effects of LEV on inflammation-related gene expression in vivo were mediated indirectly through changes in neuronal activities that affect microglia/macrophages and possibly also other types of glia. However, we cannot exclude that LEV affects genes, morphological parameters or functions of microglia/macrophages that we did not measure. LEV has been shown to affect some properties of immortalized microglial cell lines (Shima et al., 2018; Itoh et al., 2019), although it should be noted that such cell lines differ from primary microglia in many respects, including in their responses to LPS stimulation (Butovsky et al., 2014; Das et al., 2016). For the following reasons we also consider it most likely that the effects of tau reduction on inflammation-related gene expression resulted indirectly from the modulation of neuronal activities. Several lines of evidence suggest that tau reduction alters neuronal activities in ways that counteract hypersynchronous network activity (Chang et al., 2021), a type of network dysfunction that is known to change microglial activities (Eyo et al., 2017). Furthermore, neurons have robust levels of tau, whereas microglia and astrocytes express only minimal amounts of this gene product (Zhang et al., 2014; Saunders et al., 2018). However, because some oligodendrocytes express relatively high levels of tau (Zhang et al., 2014; Saunders et al., 2018), it is conceivable that tau reduction modulates interactions between oligodendrocytes and microglia, an intriguing possibility that deserves to be explored in future studies.

In general, inflammatory responses can be protective, harmful, or both (Wyss-Coray and Mucke, 2002; Meyer-Luehmann and Prinz, 2015; Sarlус and Heneka, 2017; Schartz and Tenner, 2020; Leng and Edison, 2021). It is likely that the inflammation-related gene expression changes we identified in hAPP-J20 mice enable adaptive, maladaptive, and innocent bystander responses by microglia and, possibly, other brain cells or blood-derived immune cells. In the current study, we began to dissect this range of possibility in relation to TREM2. We focused on TREM2 for four main reasons: (1) It is expressed in microglia/macrophages (refs (Efthymiou and Goate, 2017; Gratuze et al., 2018; Ulland and Colonna, 2018) and current study), (2) genetic alterations that impair the function of TREM2 strongly increase AD risk (Guerreiro et al., 2013; Jonsson et al., 2013; Efthymiou and Goate, 2017), (3) TREM2 expression was robustly increased in brain tissues of hAPP-J20 mice, and (4) this increase in TREM2 expression was effectively blocked by suppressing epilepticiform activity with LEV treatment or tau ablation. Our finding that genetic reduction of TREM2 expression enhanced both spontaneous and kainate-induced epileptiform activity strongly suggests that the increased expression of TREM2 in hAPP mice represents an adaptive microglial response to the epileptiform activity in these mice. The same may be true for TYROBP, a TREM2-related gene product whose increased expression in hAPP-J20 mice was also reduced by LEV treatment and tau ablation.

Taken together, these findings indicate that TREM2 critically contributes to microglial functions that counteract hypersynchronous network activity and, perhaps, other types of network dysfunction. They are in line with other studies implicating microglia in the regulation of neuronal activities and the suppression of epileptic network activity (Li et al., 2012; Eyo et al., 2014, 2016; Szalay et al., 2016; Badimon et al., 2020; Umpeierre and Wu, 2020; Merlìni et al., 2021). Reduced levels or hypofunction of TREM2 would be expected to increase network dysfunction, particularly under conditions that promote network hypersynchrony, such as pathologically elevated levels of hAPP metabolites or microinfarcts, which are common in AD (Corrada et al., 2016) and could lead to intermittent excitotoxicity. Microglia with reduced or deficient TREM2 may be less able to curtail aberrant neuronal activities, which could increase amyloid formation (Cirrito et al., 2005; Bero et al., 2011; Yuan and Grutzendler, 2016) and the spread of abnormal tau species (Pooler et al., 2013; Yamada et al., 2014; Wu et al., 2016). Thus, in addition to previously identified possibilities (Gratuze et al., 2018; Ulland and Colonna, 2018; Hammond et al., 2019), run-away network dysfunction is a potential mechanism by which TREM2 hypofunction may promote the development of AD. Possibly consistent with this notion, Nasu-Hakola disease, a rare genetic disorder due to complete loss of TREM2 or TYROBP function, typically causes both epilepsy and dementia (Dardiotis et al., 2017).
Although several genome-wide association studies indicate that alterations in immune-related genes contribute to AD pathogenesis, the relevant variants are rare, including those of TREM2 (Guerreiro et al., 2013; Jonsson et al., 2013; Efthymiou and Goate, 2017). The mediators that lead from disease triggers to immune dysfunction in the most common, sporadic forms of AD remain to be unraveled. Our study suggests that aberrant neuronal activity—and in particular subclinical epileptiform activity, which occurs in a substantive proportion of sporadic AD cases (Vossel et al., 2016)—could be such a mediator.

Notably, tau reduction and LEV effectively reduced network dysfunction and aberrant microglial gene expression in the presence of high hAPP/Aβ levels in old brains. These interventions might be particularly beneficial when microglia have been rendered dysfunctional by disease, as they appear to suppress aberrant neural network activity (refs. (Li et al., 2012; Eyo et al., 2014; Eyo et al., 2016; Szalay et al., 2016; Badimon et al., 2020; Umpierre and Wu, 2020; Merlini et al., 2021) and this study). Because tau reduction and LEV prevented or reversed changes in the expression of diverse inflammation-related genes, these strategies might also reduce maladaptive and pathogenic immune responses. Their effects on hAPP/Aβ-induced C1Q elevation could be a case in point, as C1Q is also increased in AD brains (Tenner et al., 2018; Schartz and Tenner, 2020) and blocking C1Q with an antibody prevents synapse loss in hAPP mice (Hong et al., 2016; Tenner et al., 2018; Schartz and Tenner, 2020) and inhibits the development of epilepsy after traumatic brain injury (Holden et al., 2021).

Limitations of the study
Our molecular profiling focused on a circumscribed set of inflammation-related transcripts and on bulk tissue from specific brain regions. It would be interesting to expand this analysis to a single-cell analysis of all genes expressed in the brain across different regions. Many biological processes are not reflected in mRNA changes because they strictly occur at the levels of proteins, lipids, and other factors. It could, therefore, be informative to extend our transcriptomics analysis to other ‘omics platforms. A limitation of most, if not all, of such studies is the inability to differentiate among adaptive, maladaptive and innocent bystander effects, an issue we addressed in regard to one of our lead hits by assessing the consequences of genetic TREM2 reduction in kainate-challenged mice. Because determining the sequence in which molecular alterations emerge can also help differentiate between causes and consequences, it could be useful to measure gene expression changes at additional ages. Experimental models simulate key aspects of a process or condition but rarely, if ever, recapitulate its full complexity. This is especially true for models of diseases as multifactorial and heterogeneous as AD. We previously validated various discoveries originally made in hAPP-J20 mice in both App knockin mice that do not overexpress APP (Orr et al., 2018; Johnson et al., 2020) and in humans with AD (Palop et al., 2003; Meilandt et al., 2008; Sanchez-Mejia et al., 2008; Verret et al., 2012; Vossel et al., 2013, 2016; Orr et al., 2015; Merlini et al., 2019). However, the hAPP-J20 model was developed to evaluate the specific pathogenic impact of FAD-mutant hAPP and, thus, does not include some other hallmarks of AD (e.g., neurofibrillary tangles). In general, it is difficult to predict whether the efficacy of therapeutic interventions observed in models designed to dissect rather than simulate the complexity of a disease will extrapolate to the human condition. It will, therefore, be important to assess the impact of LEV treatment and tau reduction also in alternate models designed to simulate the combined effects of multiple AD-related pathogenic factors. In a similar vein, while we simulated the hypofunction of TREM2 variants that increase AD risk by genetic deletion of one Trem2 allele in mice, it would be interesting to determine if expression of the actual human TREM2 variants has similar effects. Notwithstanding these caveats, our study clearly highlights that the relationship between neural network activities and microglial functions in health and disease merits further investigation and so does the extent to which reducing network dysfunctions can prevent or halt the immunopathology and clinical manifestations of AD.

STAR METHODS
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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.103245.

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AUTHOR CONTRIBUTIONS
M.D. and L.M. conceived and designed the study. M.D., W.M., E.S., S.T., G.Y., X.Y., K.H., X.W., and J.W. performed experiments. M.D., W.M., E.S., and L.M. analyzed the data. M.D. and L.M. wrote the paper with input from all coauthors.

DECLARATION OF INTERESTS
L.M. has received research funding from Cure Network Dolby Acceleration Partners (CNDAP) and has served on CNDAP’s board of managers. He has also served on the scientific advisory boards of Arvinas, Biogen, and Dolby Family Ventures and has consulted for Eisai and Sangamo Therapeutics. L.M. is a coinventor on patents held by the Gladstone Institutes that focus on tau reduction as a strategy to block neural network dysfunction.

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## STAR★METHODS

### KEY RESOURCES TABLE

| REAGENT OR RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Goat anti-rabbit IgG (H+L) cross-adsorbed secondary antibody, Alexa Fluor 594 | Invitrogen | Cat#: A-11012; RRID: AB_2534116 |
| Goat anti-rabbit IgG (H+L) cross-adsorbed secondary antibody, biotinylated | Vector Laboratories | Cat#: BA-1000-1.5 |
| Goat anti-rabbit IgG (H+L), HRP | Abcam | Cat#: ab6721 |
| Rabbit monoclonal anti-C1q | Abcam | Cat#: ab182451; RRID: AB_2732849 |
| Rabbit polyclonal anti-Aβ (3D6) | ATCC/Cell Essentials | Cat#: PTA-5130 |
| Rabbit polyclonal anti-iBA1 | Wako | Cat#: 019-19741; RRID: AB_839504 |
| Tyramide (Opal 620 fluorophore) | Akoya Biosciences | Cat#: FP1495001KT |
| **Bacterial and Virus Strains** |        |            |
| Ultrapure LPS, E. coli0111:B4 | InvivoGen | Cat#: tlr-3pelps |
| **Chemicals, Peptides, and Recombinant Proteins** |        |            |
| Kainate | Tocris | Cat#: 0222; CAS: 487-79-6 |
| Levetiracetam | ZerenexMolecular | Cat#: ZX-CF007056; CAS: 102767-28-2 |
| Thioflavin S | Sigma-Aldrich | Cat#: T1892; CAS: 1326-12-1 |
| **Critical Commercial Assays** |        |            |
| DAB substrate kit | Vector Laboratories | Cat#: SK-4100 |
| nCounter Neuroinflammation Panel, Mouse | NanoString | Cat#: 115000237 |
| RNAscope Multiplex Fluorescent Assay v2 | ACDBio | Cat#: 323100 |
| VECTASTAIN Elite ABC-HRP Kit | Vector Laboratories | Cat#: PK-6100 |
| **Deposited Data** |        |            |
| Raw and analyzed NanoString data | This study | Tables S1, S2, and S3 |
| **Experimental Models: Organisms/Strains** |        |            |
| Mouse: C57BL/6J | The Jackson Laboratory | JAX: #000664 |
| Mouse: hAPP-J20 (B6.Cg-Zbtb20^TgPDGFβ-APPswd20Lum/f2Mmjax) | Maintained in-house but available from The Jackson Laboratory | JAX: #34836 |
| Mouse: Mapt^{−/−}(B6.129X1-Mapt^{tm1Hnd/J}) | The Jackson Laboratory | JAX: #007251 |
| Mouse: Trem2^{−/−} (C57BL/6J-Trem2^{tm2Adiuj/J}) | The Jackson Laboratory | JAX: #027197 |
| **Oligonucleotides** |        |            |
| Mm00479862_g1 (Aif1) TaqMan Gene Expression Assay, FAM-MGB | ThermoFisher Scientific | Cat#: 4351370 |
| Mm00432142_m1 (C1qa) TaqMan Gene Expression Assay, FAM-MGB | ThermoFisher Scientific | Cat#: 4331182 |
| Mm03047343_m1 (C6d6) TaqMan Gene Expression Assay, FAM-MGB | ThermoFisher Scientific | Cat#: 4351370 |
| Mm01183349_m1 (Clec7a) TaqMan Gene Expression Assay, FAM-MGB | ThermoFisher Scientific | Cat#: 4351370 |
| Mm99999915_g1 (Gapdh) TaqMan Gene Expression Assay, VIC-MGB.PL | ThermoFisher Scientific | Cat#: 4448486 |
| Mm00434228_m1 (Il1b) TaqMan Gene Expression Assay, FAM-MGB | ThermoFisher Scientific | Cat#: 4351370 |
| Mm00446190_m1 (Il6) TaqMan Gene Expression Assay, FAM-MGB | ThermoFisher Scientific | Cat#: 4351370 |
| Mm04209424_g1 (Trem2) TaqMan Gene Expression Assay, FAM-MGB | ThermoFisher Scientific | Cat#: 4331182 |

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Lennart Mucke (lennart.mucke@gladstone.ucsf.edu).

Materials availability
This study did not generate unique reagents.

Data and code availability
- Tables S1, S2, and S3 include all NanoString data generated in this study. All other data reported in this paper will be shared by the lead contact upon reasonable request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the Lead Contact upon reasonable request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice
hAPP-J20 mice were generated in-house (Mucke et al., 2000). Mapt+/− mice (Stock #007251) and Trem2−/− mice (Stock #027197) were from The Jackson Laboratory. All mice were maintained on a C57BL/6J

Software and Algorithms

Cytoscape
Shannon et al. (2003)
https://cytoscape.org
RRID: SCR_00302

Fiji
Schindelin et al. (2012)
https://imagej.net/Fiji
RRID: SCR_002285

Imaris
Bitplane
https://imaris.oxinst.com
RRID: SCR_007370

JACoP (Fiji plugin)
Bolte and Cordelieres (2006)
https://imagejdocu.tudor.lu/doku.php?id=plugin:analysis:jacop_2.0:just_another_colocalization_plugin:start

LabChart Pro
ADInstruments
https://www.adinstruments.com
RRID: SCR_003420

limma (R package)
Ritchie et al. (2015)
https://bioconductor.org/packages/release/bioc/html/limma.html
RRID: SCR_010943

nSolver Analysis
NanoString
https://www.nanostring.com
RRID: SCR_003420

R
R Core Team (2014)
https://www.r-project.org/
RRID: SCR_00195

sva (R package)
Leek and Storey (2007)
https://bioconductor.org/packages/release/bioc/html/sva.html
RRID: SCR_002155

REAGENT OR RESOURCE OR SOURCE IDENTIFIER

REAGENT OR RESOURCE
Mm00449152_m1 (Tyrobp) TaqMan Gene Expression Assay, FAM-MGB
RNAscope probe Mm-C1qa
RNAscope probe Mm-Trem2-C2
RNAscope probe Mm-Tyrobp-C3

SOURCE
ThermoFisher Scientific
ACDBio
ACDBio
ACDBio

IDENTIFIER
Cat#: 4351370
Cat#: 441221
Cat#: 4041111-C2
Cat#: 408191-C3

Continued

Other
Levetiracetam formulated chow, gamma irradiated
Placebo chow, gamma irradiated

Custom made by Bio-Serv
Bio-Serv

Cat#: S7282
Cat#: S7265

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background. Experimental and control groups were sex-balanced, matched for age and background strain, and generated by the following breeding schemes: hAPP-J20 mice and WT controls were generated by breeding male heterozygous hAPP-J20 mice with female WT mice. Mapt+/– and Mapt+/- mice that did or did not express hAPP were generated by first breeding male heterozygous hAPP-J20 mice of the Mapt+/– background with female Mapt+/- mice. Experimental and control groups were then generated with F1 breeders by crossing male hAPP-J20/Mapt+/– mice with female Mapt+/- mice. To generate Trem2+/- mice and WT controls, we first bred male Trem2+/- mice with female WT mice. Experimental and control groups were then generated with F1 breeders by crossing male or female Trem2+/- mice with WT mice. Not all mice compared in individual experiments had the same parents and were raised by the same dam at the same time (littermates). Mice were housed in sex-matched groups of up to 5 mice per cage and maintained on a 12-h light/dark cycle. Unless indicated otherwise, mice were fed a regular chow diet (PicoLab Rodent Diet 5053, TestDiet and provided with water ad libitum and maintained on a 12-h light/dark cycle. Unless indicated otherwise, mice were fed a regular chow diet (PicoLab Rodent Diet 5053, TestDiet and provided with water ad libitum. At the end of experiments, mice were deeply anesthetized with an IP injection of Avertin (2,2,2-tribromomethanol, 250 mg/kg body weight) and perfused transcardially with 0.9% NaCl. For RNA analyses, hemibrains were snap frozen and maintained at −80°C. For immunohistochemical analyses, hemibrains were fixed in 4% PFA for 48 h and stored in PBS with 0.1% sodium azide at 4°C. Mice with tumors, a body condition score ≤2, or serious injuries from fighting were excluded from analyses. All animal procedures were approved by the Institutional Animal Care and Use Committee of the University of California, San Francisco.

**Drug treatments of mice**

Some mice were fed a diet containing 2,000 mg LEV per kg chow (S7282, custom made by Bio-Serv). Based on average food consumption, their daily intake of LEV was estimated at ~300 mg/kg body weight. Placebo controls were fed the same diet without LEV (S7265, Bio-Serv). LEV and placebo chow were gamma-irradiated to meet mouse husbandry requirements. At the end of the treatment period, plasma levels of LEV were determined in a randomly selected subset of LEV-treated mice (n = 12). Briefly, 20 μl of blood was taken from the left cardiac ventricle of anesthetized mice and placed on ice in EDTA-containing tubes (ThermoFisher Scientific). Blood samples were centrifuged for 10 min at 25°C and the plasma supernatant was stored at −80°C until analysis. Plasma concentrations of LEV were determined by Charles River using HPLC, tandem mass spectrometry and internal standards. Plasma LEV levels were comparable in WT mice and hAPP-J20 mice (4.63 ± 0.339 vs. 5.05 ± 0.430 μg/ml, means ± s.e.m., n = 5–7 mice/group, p = 0.486 by unpaired, two-tailed Student’s t-test). Other mice were treated with LPS or kainate. Ultrapure LPS (InvivoGen, *E. coli* 0111:B4) was dissolved in endotoxin-free water to a stock concentration of 5 mg/ml. Kainate (Millipore-Sigma) was dissolved in 0.9% saline to a stock concentration of 4 mg/ml. For IP injections, LPS and kainate were further diluted in sterile 0.9% saline. Doses are stated in the main text or figure legends.

**Primary microglial cultures**

Mixed glial cultures were prepared from WT mouse pups at postnatal day 0–1. Hippocampal and cortical tissues were microdissected and dissociated in papain (Worthington) by repeated pipetting. Cells were plated in Dulbecco’s modified Eagle’s medium (Life Technologies) with 10% fetal bovine serum and penicillin-streptomycin. Seven days after plating, flasks were agitated in an orbital shaker for 1 h and microglia were collected, counted and plated in Dulbecco’s modified Eagle’s medium containing 1% N-2 supplement (Thermo Fisher Scientific). Treatments with LEV (Zerenex Molecular) and LPS (InvivoGen, *E. coli* 0111:B4) were initiated 2 days later as described in the legend to Figure S2.

**METHOD DETAILS**

**Electroencephalography (EEG)**

Lightweight EEG plugs were constructed in-house by soldering four Teflon-coated silver wire electrodes (0.125 mm diameter) to a multichannel electrical connector. Mice were anesthetized with isoflurane, and the electrodes were implanted under the skull and over the left and right frontal cortex (+1 mm anteroposterior (AP) and ± 1 mm mediolateral (ML) relative to the bregma) and the left and right parietal cortex (−1.7 mm AP and ± 1.7 mm ML relative to the bregma). The left frontal cortex was used as the reference electrode. Mice were allowed to recover from surgery for at least 2 weeks before EEG recordings began. Digital EEG activity and videos of their locomotor activity were recorded with a PowerLab data acquisition system. All signals were acquired at a sampling rate of 2 kHz. EEG recordings were analyzed with LabChart 8 Pro software (ADInstruments). Individual epileptic spikes were detected automatically with a macro.
written in LabChart. Deflections were identified as epileptiform spikes if their amplitude was \( \geq 4 \)-fold higher than the average baseline of the trace and the absolute value of the second derivative of the slope was \( \geq 10^4 \). EEG traces and videos were also evaluated by an investigator blinded to genotype and treatment of the mice. Spikes judged to have resulted from movements or other artifacts were removed before quantitative analyses.

**Behavioral testing**

Mice were group-housed between behavioral tests. The investigators were blinded to their genotype and treatment. Behavioral tests were administered during the day time. Testing arenas were cleaned with 70% alcohol between mice. **Open field.** Spontaneous activity in an open field was assessed in a clear plastic chamber (41 \( \times \) 41 \( \times \) 30 cm) with photobeam arrays that automatically detected horizontal (16 \( \times \) 16 photobeams) and vertical (16 photobeams) movements (Flex-Field/Open Field Photobeam Activity System, San Diego Instruments). Mice were allowed to acclimate to the testing room under normal light for 1 h before testing. During testing, mice were placed in the center of a clear plastic chamber and allowed to explore the chamber for 15 min. Active place avoidance. The testing chamber consisted of a square rotating arena with a grid floor (81 \( \times \) 81 cm; Bio-Signal Group Corp.). A smaller, clear plastic circular enclosure (40 cm in diameter) with a circular lid was placed on the center of the grid floor. The arena was surrounded by large distal spatial cues. A 60° wedge within the 40 cm arena was designated as the aversive zone and was maintained in a constant position relative to the spatial cues outside the arena. During the first day, a habituation trial was conducted in which the mice were placed inside the enclosure, the arena was rotated clockwise (1 RPM), and the mouse was allowed to freely explore for 10 min with the aversive zone stimulation deactivated. On training days 1–4, mice were placed inside the enclosure and allowed to freely explore the rotating arena for 10 min with the stimulus generator activated for foot shock delivery upon entry into the aversive zone. One 10-min trial was performed per day over the course of 4 consecutive training days. 48 hours later, mice were placed inside the enclosure and allowed to freely explore the arena for 10 min with the aversive zone stimulus deactivated (probe trial). A Tracker video tracking system (Bio-Signal Group Corp.) was used to determine when the mice were inside the aversive zone. Upon entry, a 0.2 mA shock was delivered for 500 ms and repeated every 1.5 s until the mice left the aversive zone. The behavior of mice was video recorded and analyzed with Tracker (Bio-Signal Group Corp., Acton, MA). To assess learning performance in the APA, the raw number of entries into the aversive zone was transformed into rank summary scores using the Excel PERCENTRANK function, which returns the rank of a value in a dataset as a percentage of the dataset. Rank summary scores were calculated for each mouse relative to the scores of all genotypes obtained in any given trial. One rank summary score was then calculated for each mouse by averaging all of its rank summary scores across all training trials.

**NanoString neuroinflammation panel**

Total RNA was extracted from mouse hippocampi with RNeasy kits (Qiagen) and treated with DNase. For transcriptomic analysis, 0.5 μg of total RNA per sample was hybridized and multiplexed to probes from the NanoString Neuroinflammation gene expression panel (770 neuroinflammation-related mouse genes). Probes were designed and synthesized by NanoString nCounter Technologies. The target–probe complexes were captured, imaged, and digitally counted on the nCounter platform (SPRINT multiplex).

**qRT-PCR**

RNA isolated from the cortex of mice was reverse-transcribed with TaqMan reverse transcription reagents according to the manufacturer’s protocol. The resulting cDNA was analyzed by qPCR with an ABI Prism 7900 HT Sequence Detection System, TaqMan Gene Expression Master Mix, and TaqMan primers with reporter dyes. Levels of each target cDNA were detected using FAM dye. Target cDNA levels were calculated by the \( 2^{-\Delta\Delta T} \) method, using Gapdh cDNA levels in the same well (detected with VIC dye) as a reference, and expressed relative to the mean of control values.

**Histopathology**

Coronal sections of PFA-fixed hemibrains (30 μm in thickness) were prepared with a cryostat (analyses involving RNA in situ hybridization) or a freezing microtome (all others). For ThioS staining of amyloid plaques, sections were washed 3 times in PBST and incubated for 10 min in 50% ethanol and PBS containing filtered 0.0015% ThioS and washed 3 times in PBST. Sections were mounted and counterstained with TO-PRO-3 (1:5000; ThermoFisher Scientific) and cover-slipped with ProLong Diamond Antifade Mountant.
(ThermoFisher Scientific). For immunostaining of Aβ deposits, sections were incubated with 3% hydrogen peroxide and 10% methanol in PBS for 15 min at room temperature to quench endogenous peroxidase activity, and blocked for 1 h in 10% normal donkey serum, 1% non-fat dry milk, 0.2% gelatin, and PBS containing 0.5% Triton-X 100. Sections were incubated with the 3D6 primary antibody (1:1000; ATCC/Cell Essentials) diluted in 3% normal donkey serum overnight at 4°C. Sections were then washed three times in PBS with 0.25% Triton-X 100, and incubated in biotinylated anti-rabbit secondary antibody (1:500; Vector Laboratories). Signals were enhanced and visualized with Elite ABC and 3,3’-diaminobenzidine (DAB) tetrahydrochloride kits (Vector Laboratories). For immunostaining of C1Q and IBA1, free-floating sections were rinsed in PBST followed by incubation in 10 mM EDTA (pH 6.0) at 100°C for 10 min for antigen retrieval (Ghatak and Combs, 2014). After rinsing sections twice in PBS, endogenous peroxidase activity was quenched with 3% hydrogen peroxide in PBS for 10 min followed by a 1-h incubation in blocking solution containing 10% normal goat serum in PBS and 0.1% BSA. Sections were incubated in primary antibodies against C1Q (1:500; Abcam) or IBA1 (1:1000; Wako) at 4°C overnight. Sections were then washed in PBST and incubated with anti-rabbit secondary antibody conjugated to Alexa-Fluor 594 (1:500; Invitrogen) for 1 h in the dark, followed by 3 washes in PBST. For immunostaining following RNA in situ hybridization, cryosections were first hybridized with the RNA probe of interest according to the RNAscope protocol specifications. Sections were then blocked in 10% normal goat serum for 1 h and then incubated with IBA1 primary antibody at 4°C overnight. The next day, sections were washed in PBST and incubated with anti-rabbit secondary antibody conjugated to horseradish peroxidase (1:500; Abcam) for 2 h, incubated with fluorophore-labeled tyramide (1:750; Akoya Biosciences) for 10 min to intensify the signal, washed 3 times in PBST, stained with ThioS as above, and counterstained with DAPI (1:10000; ThermoFisher Scientific). Slides were imaged with an Aperio VERSA automated slide scanner using a 10× objective (for sections labeled with 3D6, anti-C1Q, or ThioS) or with an Olympus FV3000 Laser Scanning Confocal Microscope using a 20× or 40× objective (for sections labeled with ThioS, anti-IBA1, and mRNA probes).

QUANTIFICATION AND STATISTICAL ANALYSIS
Approaches to ensure robust and unbiased data collection and statistical analysis
Investigators were blinded to the genotype and treatment of mice and cultures during data collection. Group sizes were estimated based on previous results from similar experiments and from pilot studies. Statistical analyses were done with Excel (v.16.46), Prism 8.0 (GraphPad) and R (v.3.6.2). The normality of data was assessed by D’Agostino and Pearson omnibus normality test or by Shapiro-Wilk test. Variances were compared by Bartlett’s test or F-test. Differences between two groups were assessed by unpaired, two-tailed Student’s t-test and differences among multiple groups by two-way ANOVA followed by Holm-Sidak post hoc test. Null hypotheses were rejected at the 0.05 level. Statistical details for each experiment can be found in the figure legends. Additional statistical information is described in the NanoString analyses section below.

NanoString analyses
Raw files were processed with nSolver Analysis v4.0 to normalize gene expression datasets by the geNORM method (Vandesompele et al., 2002) using the geometric mean of nine housekeeping genes. The NanoString panel also included six synthetic DNA positive control targets with known concentrations and eight negative control probes designed against engineered RNA sequences not present in biological samples. Transcripts with an expression value < 20 were excluded from analysis. Expression profiles of the remaining 621 transcripts were log-transformed and quantile-normalized before further analysis. The limma package in R v3.6.2 was applied to gene expression data for differential expression analysis (Ritchie et al., 2015). T-statistics for group comparisons were calculated with robust empirical Bayes methods. p values from t-statistics were adjusted for multiple comparisons by the Benjamini–Yekuteli method (Benjamini and Yekutieli, 2001) and p < 0.05 was considered significant. For STRING analyses, gene expression changes were subjected to protein network analysis (Szklarczyk et al., 2019) using the STRING database v11.0 with a confidence score cutoff of 0.4 and allowing STRING to recommend three additional gene products of interest. The resulting protein–protein interaction networks were visualized with Cytoscape v3.7.2 (Shannon et al., 2003). To identify genes whose expression was modulated by LEV treatment and tau ablation in Figures 5K–5M, three independent groups of mice were combined in a linear model. To account for batch effects and potential hidden confounding variables, we performed a surrogate variable analysis on the expression data using the sva package in R (Leek and Storey, 2007). Estimated surrogate variables were included as covariates in the linear regression modeling analysis.
**Immunohistochemical quantification and analyses**

For the quantitative analysis of ThioS labeled plaques, three hemibrain sections per mouse were analyzed by FIJI (Schindelin et al., 2012). For each section, the region of interest was defined as the outer border of the hippocampus. ThioS positive deposits within the outlined region were thresholded and the Analyze Particles plugin was used to count the total number of plaques within the area of interest. The number of plaques in each section was divided by the area of the hippocampus. Plaques touching the border of the outline were not counted. For each plaque within the area of interest, the area occupied, circularity and mean grey values (the average pixel intensity within an area of interest) were recorded. For quantitative analysis of C1Q immunolabeled sections, three hemibrain sections per mouse were analyzed by FIJI. The area and mean grey values from the whole hippocampus were recorded. For the quantitative analysis of IBA1-and ThioS-labeled sections, three brain sections per mouse were analyzed by FIJI and Imaris. For each section, three plaques (circumference: 209 \( \mu \text{m} \pm 1.24, \text{mean} \pm \text{s.e.m} \) were chosen for analysis. To calculate the number of microglia/macrophages surrounding plaques, the Spots tool in Imaris was used to identify the number of IBA1-positive cell bodies within 25 \( \mu \text{m} \) from the perimeter of each plaque. To calculate the fraction of fluorescent signal overlap between the IBA1 and ThioS channels, Mander’s overlap coefficient (Manders et al., 1993) was calculated using the JACoP plugin in FIJI (Bolte and Cordelieres, 2006). To determine the effect of plaque distance on the soma size of microglia/macrophages, three IBA1-positive cell bodies located within 25 \( \mu \text{m} \) from the perimeter of a plaque and three IBA1-positive cell bodies located > 25 \( \mu \text{m} \) away from the nearest plaque were analyzed. Images were thresholded and the Analyze Particles plugin in FIJI was used to measure the average area and circularity of each cell body.