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Gamma frequency entrainment attenuates amyloid load and modifies microglia

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Changes in gamma oscillations (20–50 Hz) have been observed in several neurological disorders. However, the relationship between gamma oscillations and cellular pathologies is unclear. Here we show reduced, behaviourally driven gamma oscillations before the onset of plaque formation or cognitive decline in a mouse model of Alzheimer's disease. Optogenetically driving fast-spiking parvalbumin-positive (FS-PV)-interneurons at gamma (40 Hz), but not other frequencies, reduces levels of amyloid- β (A β)₁₋₄₀ and A β ₁₋₄₂ isoforms. Gene expression profiling revealed induction of genes associated with morphological transformation of microglia, and histological analysis confirmed increased microglia co-localization with A β . Subsequently, we designed a non-invasive 40 Hz light-flickering regime that reduced A β ₁₋₄₀ and A β ₁₋₄₂ levels in the visual cortex of pre-depositing mice and mitigated plaque load in aged, depositing mice. Our findings uncover a previously unappreciated function of gamma rhythms in recruiting both neuronal and glial responses to attenuate Alzheimer's-disease-associated pathology.

Activation of local circuits of excitatory and fast-spiking inhibitory neurons that resonate at 20-50 Hz gives rise to oscillations in the local field potential (LFP), called gamma oscillations¹⁻³. Although studies have demonstrated disrupted gamma in various neurological diseases, the interplay between pathology and this emergent circuit property has yet to be determined^{4,5}. In general, molecular and cellular pathology is thought to alter synaptic activity. However, in at least one disorder, Alzheimer's disease (AD), changes in synaptic activity can also feedback to alter molecular pathology. Studies have shown that increases in synaptic activity *in vivo* increase levels of $A\beta^6$, a 36- to 43-amino-acid protein, whose aggregation is thought to initiate neurotoxic events, including neuroinflammation, synaptic and neuronal loss, and tau-associated pathology⁷. We aimed to determine how gamma affects molecular pathology in a mouse model of AD. Understanding how gamma might affect disease pathogenesis has important implications for elucidating both the basic pathology of and possible therapeutic interventions for neurological diseases with altered gamma.

Reduced hippocampal gamma in 5XFAD mice

Altered gamma has been observed in multiple brain regions in several neurological and psychiatric disorders, including a reduction in spontaneous gamma synchronization in patients with AD and reduced gamma power in multiple AD mouse models^{4,5;8,9}. However, it is unclear whether gamma is altered early in disease progression and whether it affects disease pathology. Accordingly, we recorded neural activity from behaving 5XFAD mice, a well-established model of AD¹⁰. In 3-month-old mice, which have elevated levels of A β but no major plaque accumulation in the hippocampus or manifestation of learning and memory deficits¹⁰, we recorded neural activity from hippocampal subregion CA1, where gamma has been particularly well characterized

(see, for example, refs 11–14), using a virtual environment (Extended Data Fig. 1a). In CA1, gamma is present during distinct periods of activity: running, when theta oscillations (4–12 Hz) occur (Extended Data Fig. 1b, left); and quiescent behaviour, when sharp-wave ripples (SWRs) occur^{15,16} (Extended Data Fig. 1b, right). We found no clear differences in slow gamma power (20–50 Hz) between 5XFAD mice and wild-type (WT) littermates during theta (Extended Data Fig. 1c, d).

We next examined gamma during SWRs, high-frequency oscillations of 150-250 Hz lasting around 50-100 ms (Extended Data Fig. 1b, e)¹⁴. Previous work has shown that slow gamma is elevated during SWRs and that increased gamma synchrony across CA3 and CA1 during SWRs correlates with more coordinated firing between neurons¹⁶. Similarly, we found increased gamma power during SWRs (Fig. 1a, a yellow arrow indicates elevated gamma, and Extended Data Fig. 1e). The instantaneous frequencies of these slower oscillations (10-50 Hz, Methods) were a unimodal distribution centred around 40 Hz (Fig. 1b and Extended Data Fig. 1f). Comparing gamma during SWRs in WT and 5XFAD littermates, we found that gamma power was significantly lower in 5XFAD than in WT mice (Methods, Fig. 1c and Extended Data Fig. 1g, j; examples in Fig. 1c, top). Spiking was phasemodulated by gamma in both groups, although the depth of modulation was significantly smaller in 5XFAD than in WT animals (Fig. 1d and Extended Data Fig. 1h, k). Furthermore, there were fewer SWRs per unit time in non-theta periods in 5XFAD mice compared with WT (Extended Data Fig. 1i), reducing periods when gamma power is elevated (Fig. 1a, c and Extended Data Fig. 1e). These results reveal deficits in gamma modulation of CA1 spiking in a mouse model of AD before the development of major amyloid plaque accumulation or evidence of cognitive impairment. This deficit in gamma converges with evidence of gamma deficits in different mouse models of AD, and

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Figure 1 | **5XFAD** mice have reduced power in gamma during hippocampal SWRs. **a**, Average SWRs-triggered spectrograms for one mouse (left) showing gamma (yellow arrow) during SWRs (red arrow); right: frequencies below 80 Hz enlarged (n = 370 SWR). **b**, Histogram of instantaneous gamma frequencies during SWRs for mouse in **a**. **c**, Above: *z*-scored gamma power around SWR peaks for one WT and one 5XFAD mouse (mean \pm s.e.m.). Below: cumulative distribution of gamma power during SWRs (rank-sum test, n = 2,166 and 3,085 SWRs in six 5XFAD and WT mice, respectively). **d**, Above: fraction of spikes during SWRs as a function of gamma phase (mean \pm s.e.m.). Below: depth of gamma spiking modulation during SWRs (rank-sum test, bootstrap resampling, n = 2,5005XFAD and 3,000 WT phase distributions). **e**, Above: LFP trace before

reports that gamma is altered in humans with AD^{5,8,9}. Indeed, molecular deficits in Nav1.1 in humans have been linked with gamma deficits in hAPP mice⁷.

Gamma reduced $A\beta$ production in CA1

These gamma deficits during SWRs early in disease progression in this mouse model of AD prompt the question of whether gamma could affect molecular and cellular AD pathophysiology. To test this, we induced gamma optogenetically in 5XFAD/PV-Cre mice (Methods, Fig. 1e, left, and Extended Data Fig. 2a–c). We chose to drive FS-PV-interneurons at 40 Hz because we found deficits in gamma during SWRs, and instantaneous gamma frequencies during SWRs were centred around 40 Hz. Delivering 1 ms, 473 nm light pulses at 40 Hz resulted in increased power at 40 Hz in LFPs in CA1, while random stimulation did not (Fig. 1e and Extended Data Fig. 1m–o).

Aß accumulation is thought to initiate multiple neurotoxic events typical for AD pathology. Therefore, we examined whether gamma stimulation affected overall AB peptide levels in the hippocampus of 5XFAD mice. We found that 1 h of FS-PV-interneuron stimulation reduced A β_{1-40} by 53.22% and A β_{1-42} by 44.62% in the 40 Hz group, which expresses ChR2 compared with the enhanced yellow fluorescent protein (eYFP) control group, as measured in CA1 by $A\beta$ enzymelinked immunosorbent assay (ELISA) (Fig. 1f, g and raw concentration (picograms per millilitre) in Extended Data Table 1). We performed a comprehensive set of control experiments to determine whether the effect was specific to frequency, cell type, and/or rhythmicity. Neither stimulation of CamKII-positive excitatory neurons at 40 Hz nor FS-PVinterneurons at 8 Hz or random intervals significantly reduced A β levels (Fig. 1f-i and Methods). Immunohistochemical analysis using two β-amyloid-specific antibodies (Cell Signalling Technology; D54D2, BioLegend; 12F4 (ref. 17)) in CA1 confirmed these results: A β labelling and during 40 Hz optogenetic stimulation. Below: mean and standard deviation of power spectral density (n = 4 5XFAD and 3 WT mice). Red, 40 Hz; blue, random; black, no stimulation.f, Relative A β_{1-40} levels in CA1 of 5XFAD/PV-Cre mice in each stimulation condition normalized to eYFP controls (n = 8 eYFP, n = 7 40 Hz, n = 4 8 Hz, n = 6 random mice). g, As in f for A β_{1-42} (n = 4 eYFP, n = 4 40 Hz, n = 3 8 Hz, n = 3 random mice). h, Relative A β_{1-40} levels in CA1 of 5XFAD/ α CamKII-Cre mice in each stimulation condition normalized to eYFP controls (n = 6 40 Hz, n = 3 8 Hz, n = 3 random mice). i, As in h for A β_{1-42} (n = 3 mice per group). NS, not significant, *P < 0.05, **P < 0.01, ***P < 0.001 by one-way analysis of variance (ANOVA); circles indicate n, mean + s.e.m. unless otherwise specified in bar graphs.

intensity was significantly reduced by 39.5% after 40 Hz stimulation compared with eYFP controls (Fig. 2e, f, D54D2 antibody; Extended Data Fig. 2e, f, 12F4 antibody).

Brain amyloid concentration depends on A β production from amyloid precursor protein (APP) and A β clearance rates. To elucidate whether 40 Hz stimulation reduced A β production, we examined its effects on APP cleavage by measuring levels of the cleavage intermediates of APP, C-terminal fragments (CTFs) and amino (N)-terminal fragments (NTF), in the hippocampus of the 5XFAD/ PV-Cre mice. After 40 Hz stimulation, we found significantly reduced APP CTFs and NTF compared with eYFP and random controls (Fig. 2a–d and Extended Data Fig. 2d).

Previous work has shown that APP is transported and processed in recycling endosomes¹⁸, and enlarged early endosomes have been observed in brain tissue from patients with AD¹⁹. Therefore, we characterized endosomes in CA1 after stimulation using two markers, EEA1 (early endosomal antigen 1) and Rab5 (Ras-related protein encoded by the *RAB5A* gene). Altogether, the intensity of endosomal labelling of CA1 neurons significantly decreased in both EEA1 (39.7%) and Rab5 (40.1%) after 40 Hz stimulation compared with eYFP controls (Fig. 2e, g and Extended Data Fig. 2g, h). These results suggest that, in addition to observed changes in APP cleavage products, 40 Hz stimulation also alters general endosomal processing.

Gamma induced microglia morphological changes

To explore further the cellular and molecular effects of stimulation in an unbiased manner, we performed genome-wide RNA sequencing (RNA-seq) of CA1 tissue after 1 h of 40 Hz or no stimulation (eYFP) of the 5XFAD/PV-Cre mice (Fig. 3a and Extended Data Fig. 3a, b, c). Notably, 35% of all upregulated genes had their highest expression in microglia (Fig. 3b). This RNA-seq analysis strongly suggests that 40 Hz stimulation causes an alteration in the state of microglia, which

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Figure 2 | Driving 40 Hz oscillations optogenetically in hippocampus reduces A β in 5XFAD mice. a, Representative western blot showing levels of APP (CT695), APP NTF (A8967), APP CTFs (CT695), and β -actin (A5316, loading control) in CA1 of 5XFAD/PV-Cre mice expressing only eYFP or ChR2 with 40 Hz, or random stimulation conditions. One mouse per lane, two biological replicates. **b**, Relative immunoreactivity of full-length APP normalized to actin (for **b**-**d**, n = 6 mice per group). **c**, Relative immunoreactivity of APP NTF normalized to actin. **d**, Relative immunoreactivity of APP CTFs normalized to actin. **e**, Immunohistochemistry with anti-A β (D54D2, green) and anti-EEA1 (610457, red) antibodies in CA1 of 5XFAD/PV-Cre mice (scale bar, 50 µm). **f**, Relative immunoreactivity of A β normalized to eYFP controls (for **f**, **g**, n = 3 mice per group). **g**, Relative immunoreactivity of EEA1 normalized to eYFP controls. *P < 0.05, **P < 0.01, by one-way ANOVA; mean + s.e.m. in bar graphs.

is noteworthy given the accumulating evidence that microglia play a role in AD pathology²⁰. Transcriptomic changes after 40 Hz stimulation were positively correlated with changes due to increased neural activity (by NMDA (*N*-methyl-D-aspartate) and bicuculline), and negatively correlated with changes due to silencing activity (by tetrodotoxin) (Extended Data Fig. 3d). The immediate early genes *Nr4a1*, *Arc*, and *Npas4*, which are upregulated by neuronal activity, were elevated as shown by both RNA-seq and reverse transcribed quantitative PCR (RT-qPCR) (Extended Data Fig. 3e).

These transcriptomic results also suggest an engulfing state of microglia. The upregulated genes were positively correlated with gene expression changes induced by macrophage colony-stimulating factor and granulocyte macrophage colony-stimulating factor, known to promote microglial A β uptake²¹ (Extended Data Fig. 3d). RT-qPCR confirmed that upregulated genes included microglial engulfment associated genes *Cd68*, *B2m*, *Bst2*, *Icam1*, and *Lyz2* (Fig. 3c). Microglia-enriched transcriptional regulator *Irf7*, cell adhesion and migration regulator *Spp1*, and microglia proliferation markers *Csf1r* and *Csf2ra* were also upregulated (Fig. 3c). Notably, RT-qPCR showed that the expression levels of pro-inflammatory genes *Il6*, *Il1b* (*Il1-\beta*), *Itgam* (*Cd11b*), and the anti-inflammatory gene *Igf1* were not changed (Fig. 3c).

Given that 40 Hz stimulation upregulated both phagocytosis- and migration/cell adhesion-related genes, we examined morphological features of microglia activation. We used an antibody against microglial



Figure 3 | Driving 40 Hz oscillations optogenetically in hippocampus causes a distinct morphological transformation of microglia in 5XFAD mice. a, Heat map of differentially expressed genes determined by wholetranscriptome RNA-Seq of CA1 from 5XFAD/PV-Cre mice expressing only eYFP or ChR2 with 40 Hz stimulation. Normalized z-score values (high: red; low: blue) were calculated for each differentially expressed gene (row). b, Cell-type-specific expression patterns of upregulated genes after 40 Hz stimulation (MO, myelinating oligodendrocyte; OPC, oligodendrocyte progenitor cell; NFO, newly formed oligodendrocyte). c, RT-qPCR of specific upregulated genes: relative RNA levels (fold change) in CA1 of 5XFAD/PV-Cre expressing only eYFP or ChR2 with 40 Hz stimulation, normalized to eYFP controls (Student's *t*-test; n = 6mice per group). d, Immunohistochemistry with anti-Iba1 (019-19741, green) to identify microglia and anti-A β (12F4, red) antibodies in CA1 of 5XFAD/PV-Cre mice expressing only eYFP or ChR2 with 40 Hz, and random stimulation (40× objective; scale bar, 50 μ m). e, Number of Iba1-positive microglia (for **f**-**i**, one-way ANOVA; n = 4 mice per group). f, Diameter of Iba1-positive microglia cell bodies. g, Average length of Iba1-positive microglia primary processes. h, Percentage of Iba1-positive microglia cell bodies that are also A β -positive. *P < 0.05, **P < 0.01, *** $P \in 0.001$, ****P < 0.0001; mean + s.e.m. in bar graphs.

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Figure 4 | Driving 40 Hz oscillations in VC via light flicker reduces A β and amyloid plaques in 5XFAD mice. a, LFP trace in VC before and during 40 Hz light flicker (above). Power spectral density mean and s.d. (below, n = 4 5XFAD mice, five recording sessions). b, Fraction of spikes in VC over four cycles of 40 Hz flicker (left) or the equivalent time for random flicker (right, n = 4 5XFAD mice from five recording sessions, mean \pm s.e.m. across animals). For random stimulation, spiking was aligned to light turning on; grey indicates additional light-on flickers occurring randomly (Methods). c, Relative A β_{1-40} (left) and A β_{1-42} (right) levels normalized to dark, in VC of 5XFAD mice exposed to dark, light, 40 Hz, 20 Hz, 80 Hz, 40 Hz with picrotoxin (PTX), and random conditions (n = 12 dark; n = 6 light, 40 Hz, 20 Hz, 80 Hz flicker and PTX; n = 4 random mice; one-way ANOVA). d, Immunohistochemistry with anti-Iba1 (019-19741, green) and anti-A β (12F4, red) antibodies in VC of

marker Iba1 to label microglia in CA1 sections from 5XFAD/PV-Cre mice after 1 h of 40 Hz or random stimulation, or in mice expressing only eYFP (Fig. 3d). We observed almost twice as many microglia in the 40 Hz group compared with the eYFP and random control groups (Fig. 3d, e). Furthermore, microglia cell body diameter increased by 135.3% after 40 Hz stimulation compared with eYFP controls and by 138.7% compared with random stimulation (Fig. 3d, f). The lengths of microglia primary processes were reduced by 54.0% in the 40 Hz stimulation condition compared with eYFP controls and by 38.5% compared with random stimulation (Fig. 3d, g). Iba1 levels did not affect these findings, as gene expression analysis showed that Iba1 expression did not differ between conditions (Extended Data Fig. 3a, b). The increase in cell body size and decrease in process length observed after 40 Hz stimulation are consistent with a shift towards a phagocytic state of microglia²². To evaluate microglia A β uptake specifically, we measured co-localization of $A\beta$ within microglia by co-immunostaining with an Aß antibody (12F4, Methods). The percentage of microglia co-localized with $A\beta$ in the cell body increased to 85.6% after 40 Hz stimulation from 31.7% (eYFP control, Fig. 3d, h). Three-dimensional renderings of microglia further demonstrated the presence of AB within microglia (Supplementary Videos 1-3). We did not find evidence of neuronal

5XFAD mice exposed to dark or 40 Hz flicker. Right: $120 \times 200m$; arrows indicate +Iba1/+Aβ signal in cell body (scale bar, 50 µm). e, Number of Iba1-positive microglia (for e-h Student's *t*-test unpaired, n = 4 mice per group). f, Diameter of Iba1-positive microglia cell bodies. g, Average length of Iba1-positive microglia primary processes. h, Percentage of Iba1-positive microglia cell bodies that are also Aβ-positive. i, Relative Aβ₁₋₄₀ levels in VC of 6-month-old 5XFAD mice after 7 days of 1 h per day dark or 40 Hz flicker (Student's *t*-test unpaired; n = 13 mice per group). j, As in i for Aβ₁₋₄₂. k, Immunohistochemistry with anti-Aβ (D5452, green) antibody in 6-month-old VC of 5XFAD mice after 7 days of 1 h per day dark or 40 Hz flicker showing plaques (white arrows; scale bar, 50 µm). l, Number of Aβ-positive plaques; (for l, m, Student's *t*-test unpaired, n = 8 mice per group). m, Area of Aβ-positive plaques. *P < 0.05, **P < 0.01, ***P < 0.001; circles indicate n, mean + s.e.m. in bar graphs.

loss by measuring the CA1 cellular layer thickness (Extended Data Fig. 3f, g). Together, these results suggest that gamma stimulation triggers microglia to increase $A\beta$ uptake.

Light flicker entrains gamma in VC

Many studies have shown that visual stimulation can drive oscillations in the gamma range^{2,23}. In particular, flickering lights at a specific frequency can induce that frequency in primary visual cortex $(VC)^{24}$. To determine whether light flickering could entrain 40 Hz oscillations to subsequently alter A β , we exposed 5XFAD mice to 40 Hz flickering for 1 h, analogous to optogenetic stimulation that reduced A β described above (Supplementary Video 4 and Methods). In VC, we found that light flickering at 40 Hz increased power in the LFP at 40 Hz, while random interval flickering (random flicker) and dark exposure did not (Fig. 4a and Extended Data Fig. 4a). All induced similar firing rates (Extended Data Fig. 4b, c). Spiking increased and decreased concomitantly as the light flickered on and off, resulting in spiking entrained to 40 Hz during 40 Hz flicker but not during random flicker (Fig. 4b). Recordings from saline above the brain exhibited no change in 40 Hz power during 40 Hz flicker, showing that this effect was not due to photoelectric effects or electrical noise (Extended Data Fig. 4d, e).

Light flicker reduces $A\beta$ levels in VC

Given that 40 Hz light flicker entrains 40 Hz oscillations in VC, we aimed to determine whether 40 Hz flicker could reduce A β levels. Three-month-old 5XFAD mice were placed in a dark box and exposed to either light flicker at different frequencies (20, 40, or 80 Hz), random flicker, constant light-on (light), or dark for 1 h. One hour after 1 h of 40 Hz flicker, we observed that $A\beta_{1-40}$ levels in VC were reduced by 57.96% and A β_{1-42} levels by 57.97% compared with dark controls (as measured by AB ELISA, Fig. 4c). The effect was specific to 40 Hz flicker as neither constant light nor 20 Hz, 80 Hz, or random flicker significantly reduced A β levels compared with dark and light controls (Fig. 4c). We also found no change in A^β levels in somatosensory barrel cortex and hippocampus after 40 Hz flicker (Extended Data Fig. 5a-h). When we pre-treated 5XFAD mice with a low-dose of GABA_A antagonist (picrotoxin, 0.18 mg/kg (ref. 25)), the effects of 40 Hz flicker on A^β levels were completely abrogated, indicating that GABAergic neurotransmission is necessary for this effect (Fig. 4c). To demonstrate that these effects extend beyond the 5XFAD mouse, we examined the effect of 40 Hz flicker in APP/PS1 mice, another wellvalidated AD model²⁶, and found significantly reduced $A\beta_{1-40}$, by 20.80%, and a non-significant trend of reduced A β_{1-42} by 37.68% (Extended Data Fig. 6a). Furthermore, in 9-month-old WT mice, we found a 58.2% reduction in endogenous mouse $A\beta_{1-40}$ after 1 h of 40 Hz flicker (Extended Data Fig. 6b). The reduction of endogenous mouse $A\beta_{1-40}$ in WT animals reveals that these effects are not restricted to transgenic APP expression or mutant APP; rather, they extend to $A\beta$ produced from endogenous APP driven by its endogenous promoter.

Next, we investigated whether 40 Hz flicker alters microglia activity in VC in a similar manner to the way 40 Hz optogenetic stimulation altered CA1 microglia. While microglia number was unchanged (Fig. 4d, e), microglia cell body diameter increased by 165.8% after 40 Hz flicker compared with dark controls (Fig. 4d, f). Microglia primary process lengths were reduced by 37.7% after 40 Hz flicker compared with dark controls (Fig. 4d, g). Consistent with this morphology, which indicates enhanced engulfment activity²², Aβ/Iba1 co-localization in the cell body was increased to 90.8% after 40 Hz flicker from 57.3% in the dark condition, indicating more Aβ-bearing microglia (Fig. 4d, h, P < 0.01). To better resolve the morphological change in microglia, we used CLARITY²⁷ to create three-dimensional renderings of microglia from 100 µm sections of VC (Supplementary Videos 5, 6). To demonstrate that microglia indeed engulf A β in the 5XFAD mouse, we purified microglia from 5XFAD and WT animals using fluorescenceactivated cell sorting (FACS) and analysed A β levels via human ELISA. We found that microglia-specific levels of $A\beta$ are significantly higher in 5XFAD animals than in WT controls (Extended Data Fig. 7a, b). Synaptophysin levels did not change between dark and 40 Hz flicker conditions, indicating that microglia activation probably did not affect synapse number (Extended Data Fig. 7c, d; NS, not significant, n = 4 mice). Thus, 40 Hz oscillations induced non-invasively via sensory entrainment reduced AB abundance and promoted microglia/ A β interactions.

Light flicker decreases plaque load in VC

We next assessed whether 40 Hz flicker was effective in treating animals that have amyloid plaques. Because the effects of 40 Hz flicker on soluble A β levels were transient, lasting 12 h but not 24 h (Extended Data Fig. 8a), we hypothesized that we would need repeated exposure to affect insoluble A β . Thus, we treated 6-month-old 5XFAD mice, which have amyloid plaque pathology in many brain regions including VC¹⁰, for 1 h daily over 7 days with 40 Hz flicker or dark, and analysed VC tissue 24 h later. We found that 7 days of 1 h 40 Hz flicker reduced both soluble A β_{1-40} and A β_{1-42} levels, by 60.5% and 51.7% respectively, (Fig. 4i, j) and insoluble A β_{1-40} and A β_{1-42} levels by 43.7% and 57.9% respectively (Fig. 4i, j). Immunohistochemical analysis showed that 40 Hz flicker significantly reduced plaque number in VC by 67.2% compared with dark controls (Fig. 4k, l) and plaque size by 63.7%

(Fig. 4k, m). Taken together, these experiments identify a non-invasive treatment with a profound effect on amyloid plaque pathology. We next determined if 40 Hz flicker reduced tau phosphorylation by immunohistochemistry, another AD-related pathology. Using the TauP301S tauopathy mouse model²⁸ we found that 7 days of 1 h 40 Hz visual flicker treatment reduced phosphorylated tau serine202 and serine404/ threonine403/serine400 levels in VC by 41.2% and 42.3%, respectively, and triggered microglia responses similar to those observed in 5XFAD mice (Extended Data Fig. 9a–k).

Discussion

Gamma oscillations are thought to be important for higher cognitive functions and sensory responses^{2,12,23}. Here, we demonstrated that entraining oscillations and spiking at 40 Hz, using optogenetics in the hippocampus of 5XFAD mice and using a non-invasive light flicker treatment to affect primary VC in multiple mouse models, resulted in a marked reduction of A β peptides. We also found a concomitant microglia response after 40 Hz entrainment.

The robust reduction of total amyloid levels was probably mediated both by decreased amyloidogenesis and by increased amyloid endocytosis by microglia. Thus, it appears that driving 40 Hz gamma oscillations may induce an overall neuroprotective response that recruits both neurons and microglia. The fact that GABA_A antagonist treatment completely abrogated the effects of 40 Hz stimulation on A β levels strongly suggests that GABAergic neurotransmission is critical for these effects.

Flicker stimulation at 40 Hz reduced A β in multiple mouse models, including 5XFAD, APP/PS1, and WT mice. This replication in multiple mouse models shows that these findings are not specific to one animal model and, importantly, extend to situations where A β is produced from APP expressed by its physiological promoter as it is in WT animals. In addition, we found that 40 Hz oscillations reduced phosphorylated tau staining in a mouse model of tauopathy, TauP301S, showing that the protective effects of gamma stimulation generalize to other pathogenic proteins.

These observations indicate that entraining gamma oscillations may provide a broad spectrum of systemic effects in the brain, including in non-neuronal cells, to attenuate AD-related pathology. Because this approach is fundamentally different from previous AD therapies²⁹, further study is needed to determine whether it will be therapeutic in human AD.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Supplementary Information is available in the online version of the paper.

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Author Information RNA-seq data available at Gene Expression Omnibus under accession number GSE77471. Other data are publicly available upon request. Reprints and permissions information is available at www.nature.com/reprints. The authors declare competing financial interests: details are available in the online version of the paper. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to L.-H.T. (Ihtsai@mit.edu).

METHODS

Animals. All animal work was approved by the Committee for Animal Care of the Division of Comparative Medicine at the Massachusetts Institute of Technology. Adult (3-month-old) male double transgenic 5XFAD Cre mice were produced by crossing 5XFAD transgenic mice with the transgenic PV or CW2 promoter driven Cre line. Adult (5-month-old) male and female APP/PS1 mice were gifted from the Tonegawa laboratory. Adult (4-month-old) male TauP301S mice were obtained from the Jackson Laboratory. Nine-month-old WT mice (C57Bl/6) were obtained from the Jackson Laboratory. Mice were housed in groups of three to five on a standard 12h light/12h dark cycle, and all experiments were performed during the light cycle. Food and water were provided *ad libitum* unless otherwise noted. Littermates were randomly assigned to each condition by the experimenter. The experimenter was blind to animal genotypes during tissue processing and electrophysiological recording and analysis. No animals were excluded from analysis. Adeno-associated viral vectors. Adeno-associated viral (AAV) particles of serotype 5 were obtained from the Vector Core Facility at The University of North Carolina at Chapel Hill. The AAV5 virus contained a channelrhodopsin-2 (ChR2) fused to eYFP in a double-floxed, inverted, open-reading-frame (DIO) driven by the EF1 a promoter (Extended Data Fig. 2a). An AAV-DIO-eYFP construct was used as a control.

Surgical procedures. Three-month-old 5XFAD/PV-Cre or CW2 mice were an aesthetized with an intraperitoneal (i.p.) injection of a mixture of ketamine (1.1 mg/kg) and xylazine (0.16 mg/kg). A small craniotomy was made 2.0 mm posterior to bregma and 1.8 mm lateral to the midline on the left side. Virus was delivered through a small durotomy by a glass micropipette attached to a Quintessential Stereotaxic Injector (Stoelting). The glass micropipette was lowered to 1.2 mm below the brain surface. A bolus of 1 µL of virus (AAV-DIO-ChR2–eYFP or AAV-DIO–eYFP; 2×10^{12} viral molecules per millilire) was injected into the CA1 region of the hippocampus at 0.075 µL min⁻¹. The pipette remained in place for 5 min following the injection before being retracted from the brain. A unilateral optical fibre implant (300µm core diameter; Thor Labs) was lowered to 0.9 mm below the brain surface about the injection site. Two small screws anchored at the anterior and posterior edges of the surgical site were bound with dental glue to secure the implant in place.

For electrophysiological recordings, adult (3-month-old) male 5XFAD/PV-Cre and 5XFAD negative littermates (for CA1 recordings), or 5XFAD and their WT littermates (for VC recordings) mice were anaesthetized using isoflurane and placed in a stereotactic frame. The scalp was shaved, ophthalmic ointment (Puralube Vet Ointment, Dechra) was applied to the eyes, and Betadine and 70% ethanol were used to sterilize the surgical area. For CA1 recordings, a craniotomy (in millimetres, from bregma: -2 anterior/posterior, 1.8 medial/lateral) was opened to deliver $1\,\mu\text{L}$ of virus to CA1 (as described above). The target craniotomy site for LFP recordings was marked on the skull (in mm, from bregma: -3.23 anterior/ posterior, 0.98 medial/lateral for CA1 and 2.8 anterior/posterior, 2.5 medial/lateral for VC), three self-tapping screws (F000CE094, Morris Precision Screws and Parts) were attached to the skull, and a custom stainless steel headplate was affixed using dental cement (C&B Metabond, Parkell). On the day of the first recording session, a dental drill was used to open the LFP craniotomies (300-400 µm diameter) by first thinning the skull until \sim 100 μ m thick, and then using a 30-gauge needle to make a small aperture. The craniotomy was then sealed with a sterile silicone elastomer (Kwik-Sil WPI) until recording that day and in between recording sessions.

Optogenetic stimulation protocol. Two to four weeks after virus injection and implant placement (which provided time for the mice to recover and undergo behaviour training for animals used for electrophysiology, and the virus to express in the neurons), CA1 neurons were optogenetically manipulated. A 200 mW, 4,793 nm DPSS laser was connected to a patch cord with a fibre channel/physical contact connector at each end. During the experiment, 1 mW (measured from the end of the fibre) of optical stimulation was delivered for 1 h. For molecular and biochemical analyses, each animal received one of three stimulation protocols: 8 Hz, 40 Hz, or random stimulation (light pulses were delivered with a random interval determined by a Poisson process with an average frequency of 40 Hz). eYFP control animals received 40 Hz stimulation. For electrophysiological recordings, each animal received all stimulation conditions interleaved during recordings. Visual stimulation protocol. Fifteen minutes before the experiment, 5XFAD mice were treated with saline (control) or picrotoxin (0.18 mg/kg)²⁵. For molecular and biochemical analyses, mice were then placed in a dark chamber illuminated by a light-emitting diode (LED) bulb and exposed to one of five stimulation conditions: dark, light, 20 Hz, 40 Hz (12.5 ms light on, 12.5 ms light off, 60 W), 80 Hz flicker for 1h. For electrophysiological recordings, each animal received dark, light, 40 Hz flicker, or random (light pulses were delivered with a random interval determined by a Poisson process with an average interval of 40 Hz) stimulation conditions interleaved in 10s blocks during recordings.

Behaviour training and virtual reality environment (VR) for electrophysiology. For CA1 recordings, head-fixed animals ran on an 8-inch spherical treadmill supported by an air cushion through a virtual reality environment, as described in ref. 30. The motion of the spherical treadmill was measured by an optical mouse and fed into virtual reality software³¹, running in MATLAB (version 2013b, Mathworks). The virtual environment consisted of a linear track with two small enclosures at the ends where the animal could turn (Extended Data Fig. 1a). Animals were rewarded with sweetened condensed milk (diluted 1:2 in water) at each end of the track for alternating visits to each end of the track.

Animals learned to run on the virtual linear track over approximately 1 week. The animals were left to recover from the surgery for 1 week, and habituated to handling for 1–2 days before behavioural training began. To learn to manoeuvre on the treadmill and get comfortable in the testing environment, on the first 2 days of training the animals were placed on the spherical treadmill with the virtual reality system off and were rewarded with undiluted sweetened condensed milk. On the second day of training on the spherical treadmill, the animals' food was restricted to motivate them to run. Animals were restricted to no more than 85% of their baseline weight and typically weighed over 88% of their baseline weight. From the third day until the end of training (typically 5–7 days) the animals were placed on the treadmill for increasing amounts of time (30 min to 2 h) running in the VR linear track. Animals were rewarded with diluted (1:2) sweetened condensed milk at the end of the linear track after traversing the length of the track. Between recording sessions, animals were given refresher training sessions to maintain behavioural performance.

For VC recordings, animals ran on the spherical treadmill while exposed to dark, light, or light-flickering conditions (described below in data acquisition). Before recordings, animals learned to manoeuvre on the treadmill and get comfortable in the testing environment by being placed on the spherical treadmill (with the virtual reality system off) and receiving a reward of undiluted sweetened condensed milk.

Electrophysiology data acquisition. For optogenetic stimulation of CA1 during recording, a 300 μ m core optical fibre was advanced through the craniotomy used to deliver virus to CA1 to a depth of 900 μ m into the brain. Light pulses that were 1 ms and 1 mW (measured from the end of the fibre) were delivered via a 473 nm DPSS (diode pumped solid state) laser (as described above).

To avoid photoelectric artefacts, neural activity was recorded with glass electrodes. LFP electrodes were pulled from borosilicate glass pipettes (Warner) on a filament-based micropipette puller (Flaming-Brown P97, Sutter Instruments), to a fine tip, which was then manually broken back to a diameter of ${\sim}10{-}20\,\mu\text{m}$ and filled with sterile saline. For CA1 recordings the LFP electrode was advanced through the LFP recording craniotomy at an angle 60 degrees posterior to the coronal plane and 45° inferior to the horizontal plane until clear electrophysiological signatures of the hippocampal stratum pyramidale layer were observed (${\sim}600{-}1000\,\mu\text{V}$ theta waves while the animal was running, clearly distinguishable SWRs during immobility, and multiple spikes greater than 150 μ V; Extended Data Fig. 1b). For VC recordings, the LFP electrode was advanced vertically through the LFP recording craniotomy to a depth of 600–900 μm and multiple spikes greater than 150 μ V were observed.

Data were acquired with a sampling rate of 20 kHz and bandpass filtered 1 Hz to 1 kHz. Animals ran on the spherical treadmill or rested for prolonged periods. For optogenetic simulation sessions, data were recorded for 30 min before any stimulation began. Then stimulation was delivered at gamma (40 Hz), random (as described under Optogenetic stimulation protocol), or theta (8Hz) frequency for 10s periods interleaved with 10s baseline periods (no stimulation). In two animals, stimulation of each type or baseline was delivered for 5 min periods instead of 10s periods. Each 30 min of stimulation recordings were followed by 5-30 min of recording with no stimulation. For visual light flicker simulation sessions, LED striplights surrounding the animal lights were flickered at gamma (40 Hz), random (described above in Visual stimulation protocol), theta (8 Hz), or 20 Hz frequency for 10 s periods, or were on continuously for 10 s periods, interleaved with 10 s periods with lights off. A few recordings were made above the brain surface during light flicker to ensure that the lights did not create electrical or photoelectric noise during recording. Recording sessions were terminated after approximately 3-5h. Animals were 3-4 months old at the time of recording.

Spike detection. Spikes were detected by thresholding the 300–6,000 Hz band-passed signal. Threshold was the median of the filtered signal plus five times a robust estimator of the standard deviation of the filtered signal (median/0.675) to avoid contamination of the standard deviation measure by spikes³².

LFP. Recorded traces were downsampled to $2\,\rm kHz$ and then bandpass filtered between 1 and 300 Hz.

Theta and SWR detection. Activity across the hippocampal network changes markedly when animals run or sit quietly, and these changes are often referred to

as different network states. These network states are clearly distinguishable by the presence or absence of LFP oscillations in different frequency bands^{12,13}. When animals ran, we observed large theta (4–12 Hz) oscillations in CA1 as others have shown (Extended Data Fig. 1b, left)^{13,30,33,34}. When animals sat quietly, theta oscillations were no longer visible and we recorded SWRs, high-frequency oscillations of 150–250 Hz that last around 50–100 ms and are associated with bursts of population activity, as others have observed (Extended Data Fig. 1b, right)^{15,16}.

SWRs were detected (Fig. 1a-d and Extended Data Fig. 1d-i) when the envelope amplitude of the filtered trace was greater than four standard deviations above the mean for at least 15 ms. The envelope amplitude was calculated by taking the absolute value of the Hilbert transform of the filtered LFP. We also confirmed our results held when using a higher threshold for SWR detection, six standard deviations above the mean, which detects larger SWRs (Extended Data Fig. 1j, k). To detect theta (Extended Data Fig. 1c, d), the LFP was bandpass filtered for theta (4-12 Hz), delta (1-4 Hz), and beta (12-30 Hz) using an FIR equiripple filter. The ratio of theta to delta and beta ('theta ratio') was computed as the theta envelope amplitude divided by the sum of the delta and beta envelope amplitudes. Theta periods were classified as such when the theta ratio was greater than one standard deviation above mean for at least 2s and the ratio reached a peak of at least two standard deviations above mean. Non-theta periods were classified as such when the theta ratio was less than one for at least 2 s. SWRs, theta periods, and non-theta periods were visually inspected to ensure that these criteria accurately detected SWRs, theta periods, and non-theta periods, respectively.

Power spectrum. Spectral analysis was performing using multitaper methods (Chronux toolbox, time-bandwidth product = 3, number of tapers = 5). For examining power spectra without stimulation (Extended Data Fig. 1c, d), only theta periods were included: theta periods greater than 5 s long were divided into 5 s trials and the average power spectral density was computed for each animal over these trials. For examining power spectra during optogenetic (Fig. 1e and Extended Data Fig. 1l) and visual stimulation (Fig. 4a and Extended Data Fig. 4a), data were divided into 10 s trials of each stimulation condition or baseline periods, and the average power spectral density was computed for each animal over these trials.

Gamma during SWRs. Spectrograms were computed using multitaper methods (Chronux toolbox). The spectrogram was computed for each SWR including a window of 400 ms before and after the peak of the SWR. Then a z-scored spectrogram was computed in each frequency band using the mean and standard deviation of the spectrogram computed across the entire recording session to create a normalized measure of power in units of standard deviation (Fig. 1a and Extended Data Fig. 1e). Instantaneous frequency of gamma during SWRs was computed by bandpass filtering the LFP for 10-50 Hz, taking the Hilbert transform, then taking the reciprocal of the difference in peaks of the transformed signal (Fig. 1b and Extended Data Fig. 1f). Gamma power before, during, and after SWRs was computed by filtering the LFP for low gamma (20-50 Hz) and taking the amplitude of the envelope of the Hilbert transform to get the mean gamma power in 100 ms bins centred on the SWR peak. This was normalized by the mean and standard deviation of the amplitude of the envelope for the entire recording session to get z-scored gamma power for each bin around each SWRs (Fig. 1c and Extended Data Fig. 1g, j). Phase modulation by gamma during SWRs was computed by bandpass filtering the LFP for gamma (20-50 Hz), taking the Hilbert transform, and determining the phase of the resulting signal for each spike that occurred during SWRs (Extended Data Fig. 1h). To measure differences in phase modulation between 5XFAD and WT animals, we used resampling with replacement: a subset of 100 spikes from each recording was randomly selected to create a phase modulation distribution and this was repeated 500 times for each recording (Fig. 1d and Extended Data Fig. 1k). We then measured the depth of modulation for the spike-gamma phase distribution by computing the difference between the peak and trough divided by the sum of the peak and trough for each distribution (Fig. 1d and Extended Data Fig. 1k).

Differences in firing during stimulation. To plot stimulus-evoked multiunit firing histograms, spikes were binned in 2.5 ms bins for 100 ms after the start of each light-on pulse and the fraction of spikes in each bin was computed. Mean and standard error were then computed across all light-on periods. To compute differences in multi-unit firing rate between conditions, firing rates were computed for each 10 s period of stimulation or baseline (total number of spikes divided by duration of period). Differences in firing rate were taken between nearby periods of the relevant type of stimulation (firing rate in gamma stimulation period minus baseline or random periods for optogenetic stimulation, firing rate in gamma stimulation). Differences from all animals were plotted in histograms (Extended Data Figs 1m and 4c) and the median and quartiles of the multiunit firing rates per 40 Hz stimulation, random stimulation, and no stimulation period for each animal were plotted in box plots (Extended Data Figs 10 and 4d).

Immunohistochemistry. Mice were perfused with 4% paraformaldehyde under deep anaesthesia, and the brains were post-fixed overnight in 4% paraformaldehyde. Brains were sectioned at $40\,\mu m$ using a vibratome (Leica). Sections were permeabilized and blocked in PBS containing 0.2% Triton X-100 and 10% normal donkey serum at room temperature for 1 h. Sections were incubated overnight at 4°C in primary antibody in PBS with 0.2% Triton X-100 and 10% normal donkey serum. Primary antibodies were anti-EEA1 (BD Transduction Laboratories; 641057), anti-β-amyloid (Cell Signaling Technology; D54D2), anti-Iba1 (Wako Chemicals; 019-19741), anti-parvalbumin (Abcam; ab32895), and anti-Rab5 (Enzo Life Sciences; ADI-KAP-GP006-E). To confirm ELISA experiments, the anti-AB antibody D54D2 was used because it allowed for co-labelling with EEA1 and the anti-A β antibody 12F4 was used because it does not react with APP, allowing us to determine whether our labelling was specific to A^β. For co-labelling experiments, the anti-A β antibody 12F4 (Biolegend; 805501) was used. Primary antibodies were visualized with Alexa-Fluor 488 and Alex-Fluor 647 secondary antibodies (Molecular Probes), and cell nuclei visualized with Hoechst 33342 (Sigma-Aldrich; 94403). Images were acquired using a confocal microscope (LSM 710; Zeiss) with a 40× objective at identical settings for all conditions. Images were quantified using ImageJ 1.42q by an experimenter blind to treatment groups. For each experimental condition, two coronal sections from at least three animals were used for quantification. Scale bars are $50\,\mu\text{m}.$ For CA1 imaging, the analysis was restricted to the pyramidal cell layer, except in the case of Iba1+ cells analysis, where the whole field of view was required to image an adequate number of cells. ImageJ was used to measure the diameter of Iba1+ cell bodies and to trace the processes for length measurement. In addition, the Coloc2 plugin was used to measure co-localization of Iba1 and A_β. Imarisx64 8.1.2 (Bitplane, Zurich, Switzerland) was used for three-dimensional rendering. For counting the 'plaque number', deposits of at least 10µm were included.

CLARITY. Fixed brains were sliced into 100 µm coronal sections on a vibratome (Leica VT100S) in 1× PBS. Sections containing VC were selected, with reference to the Allen Mouse Brain Atlas, and incubated in clearing buffer (pH 8.5-9.0, 200 mM sodium dodecylsulfate, 20 mM lithium hydroxide monohydrate, 4 mM boric acid in double-distilled H2O) for 2 h, shaking at 55 °C. Cleared sections were washed 3×10 mins in $1 \times PBST$ (0.1% Triton-X100/1XPBS) and put into blocking solution (2% bovine serum albumin/1× PBST) overnight, shaking at room temperature²⁷. Subsequently, three 1 h washes in $1 \times PBST$ were performed, shaking at room temperature. Sections were then incubated at 4 °C for 2 days, shaking, with anti-\beta-amyloid (Biolegend; 805501) and anti-Iba1 (Wako Chemicals; 019-19741) primary antibodies, diluted to 1:100 in $1 \times$ PBST. Another set of 3×1 h washes in $1 \times$ PBST was conducted before sections were incubated for 9h, shaking at room temperature, in 1:100 1× PBS-diluted secondary antibody mixture. Fragmented Donkey Anti-Rabbit Alexa Fluor 488 (Abcam; ab175694) and Anti-Mouse 568 (Abcam; ab150101) secondary antibodies were used to visualize the primary antibody labelling. Halfway through this incubation period, Hoechst 33258 (Sigma-Aldrich; 94403) was spiked into each sample at a 1:250 final dilution. Sections were then washed overnight in $1 \times PBS$, shaking at room temperature. Before mounting for imaging, slices were incubated in refractive index matching solution (75 g Histodenz, 20 mL 0.1 M phosphate buffer, 60 mL double-distilled H₂O) for 1 h, shaking at room temperature. Tissue sections were mounted onto microscopy slides with coverslips (VWR VistaVision, VWR International, LLC, Radnor, Pennsylvania, USA) using Fluromount G Mounting Medium (Electron Microscopy Sciences, Hatfield, Pennsylvania, USA). Images were acquired on a Zeiss LSM 880 microscope with the accompanying Zen Black 2.1 software (Carl Zeiss Microscopy, Jena, Germany). Section overview and cellular-level images used for threedimensional reconstruction were taken using a Plan-Apochromat 63×/1.4 oil differential interference contrast objective. Imarisx64 8.1.2 (Bitplane, Zurich, Switzerland) was used for three-dimensional rendering and analysis.

Western blot. CA1 whole-cell lysates were prepared using tissue from 3-month-old male 5XFAD/PV-Cre mice. Tissue was homogenized in 1 ml RIPA (50 mM Tris HCl pH 8.0, 150 mM NaCl, 1% NP-40, 0.5% sodium deoxycholate, 0.1% SDS) buffer with a hand homogenizer (Sigma), incubated on ice for 15 min, and rotated at 4 °C for 30 min. Cell debris was isolated and discarded by centrifugation at 14,000 r.p.m. for 10 min. Lysates were quantitated using a nanodrop, and 25 μ g protein was loaded on a 10% acrylamide gels. Protein was transferred from acrylamide gels to PVDF membranes (Invitrogen) at 100 V for 120 min. Membranes were blocked using bovine serum albumin (5% w/v) diluted in TBS:Tween. Membranes were incubated in primary antibodies overnight at 4°C and secondary antibodies at room temperature for 90 min. Primary antibodies were anti-APP (Invitrogen; PAD CT695), anti-APP (Sigma; A8967), and anti-β-actin (Abcam; ab9485). Secondary antibodies were quantified using ImageJ 1.46a and normalized to values of β-actin. We examined tau protein solubility using sequential protein extraction as described in

ref. 28. We then probed the detergent insoluble tau fraction using an antibody against Tau5 (Thermo Fisher Scientific; AHB0042).

ELISA. CA1 or VC was isolated from male mice, lysed with PBS or 5M Guanidine HCl, and subjected to A β measurement with the use of mouse (for WT experiments) or human (for all other experiments) A β_{1-40} or A β_{1-42} ELISA kit (Invitrogen) according to the manufacturer's instructions. We lysed the tissue in phosphate-buffered saline (PBS) to extract the PBS soluble A β fraction. The soluble A β fraction probably contained monomeric and oligomeric A β . Tissue was further treated with guanidine HCl to extract the insoluble A β fraction. A β_{1-42} was below detectable levels for both flicker and control groups in WT VC and microglia-specific samples.

Genome-wide RNA-seq. Total RNA was extracted from CA1 isolates using the RNeasy kit (Qiagen). Purified mRNA was used for RNA-seq library preparation using the BIOO NEXTflex kit (BIOO 5138-08) according to the manufacturer's instructions. Total mRNA (1 μg) was subject to a sequential workflow of poly-A purification, fragmentation, first strand and second strand synthesis, DNA end-adenylation, and adaptor ligation. The libraries were enriched by 15 cycles of PCR reactions and cleaned with Agencourt AMPure XP magnetic beads (Beckman Coulter). The quality of the libraries was assessed using an Advanced Analyticalfragment Analyzer. The bar-coded libraries were equally mixed for sequencing in a single lane on the Illumina HiSeq 2000 platform at the MIT BioMicro Center. The raw fastq data of 50-bp single-end sequencing reads were aligned to the mouse mm9 reference genome using TopHat2.0. The mapped reads were processed by Cufflinks 2.2 with UCSC mm9 reference gene annotation to estimate transcript abundances, and test for differential expression. An average of 26,518,345 sequencing reads was obtained from three stimulated and three non-stimulated mice. Relative abundance of transcript was measured by fragments per kilobase of exon per million fragments mapped (FPKM). Gene differential expression test between treated and untreated groups was performed using Cuffdiff module with an adjusted *P* value <0.05 for statistical significance (GEO accession number GSE77471).

To understand the cellular and molecular mechanisms from our RNA-seq data, 14 of publicly available RNA-seq datasets³⁵ were processed for cell-type-specific analysis. Additionally, 60 publicly available neuron-, microglia-, and macrophagespecific RNA-seq datasets under different chemical and genetic perturbations³⁶⁻⁴¹ were downloaded and processed using TopHat/Cufflinks pipeline for gene set enrichment (GSEA) statistical analysis. GSEA was used to determine whether a defined gene set from our RNA-seq data are significantly enriched at either direction of a ranked gene list from a particular perturbation study. Genes detected in the public RNA-seq datasets were ranked by log₂ values of fold change (case versus control), from positive to negative values. A defined gene set (in our case, up- or downregulated genes upon gamma treatment) was considered significantly correlated with a perturbation-induced transcriptomic change (either up- or downregulation) when both nominal P value and false discovery rate q value were less than 0.05. The sign of the calculated normalized enrichment score (NES) indicates whether the gene set is enriched at the top or the bottom of the ranked list. The heatmap for differentially expressed genes was generated using a custom R script, and z-score values across all libraries for each gene were calculated on the basis of the gene FPKM values. The box plots for cell-type specificity analysis were also generated by the R program, on the basis of gene FPKM values.

Quantitative RT–PCR. The CA1 subregion was isolated from hippocampus of 3-month-old male 5XFAD/PV-Cre mice. Tissue was rapidly frozen using liquid nitrogen and stored at -80 °C, and RNA extracted using the RNeasy kit according to the manufacturer's protocol (Qiagen). RNA (3µg) was treated with DNase I (4 U, Worthington Biochemical Corporation), purified using RNA Clean and Concentrator-5 Kit (Zymo Research) according to the manufacturers' instructions, and eluted with 14µL DEPC-treated water. For each sample, 1µg RNA was reverse transcribed in a 20µL reaction volume containing random hexamer mix and Superscript III reverse transcriptase (50 U, Invitrogen) at 50 °C for 1 h. First strand cDNAs were diluted 1:10 and 1µL were used for RT-qPCR amplification in a 20µL reaction (SsoFast EvaGreen Supermix, Bio-Rad) containing primers (0.2µM). Relative changes in gene expression were assessed using the $2^{-\Delta\Delta Ct}$ method.

Isolation of microglia from VC. The primary VC (V1 region) was rapidly dissected and placed in ice-cold Hanks' balanced salt solution (HBSS) (Gibco by

Life Technologies, catalogue number 14175-095). The tissue was then enzymatically digested using the Neural Tissue Dissociation Kit (P) (Miltenyi Biotec, catalogue number 130-092-628) according to the manufacturer's protocol, with minor modifications. Specifically, the tissue was enzymatically digested at 37 °C for 15 min instead of 35 min and the resulting cell suspension was passed through a $40 \,\mu\text{m}$ cell strainer (Falcon Cell Strainers, Sterile, Corning, product 352340) instead of a MACS SmartStrainer, $70 \,\mu\text{m}$. The resulting cell suspension was then stained using allophycocyanin (APC)-conjugated CD11b mouse clone M1/70.15.11.5 (Miltenyi Biotec, 130-098-088) and phycoerythrin (PE)-conjugated CD45 antibody (BD Pharmingen, 553081) according to the manufacturer's (Miltenyi Biotec) recommendations. FACS was then used to purify CD11b and CD45 positive microglial cells. The cells were sorted directly into 1 × PBS (Extended Data Fig. 6a).

Code availability. Code is publicly available upon request from the corresponding author.

Statistics. For electrophysiological data that were not normally distributed, results are presented as medians and quartiles unless otherwise noted. Two-sided Wilcoxon rank sum tests for equal medians were performed to determine whether distributions were significantly different, and Wilcoxon signed rank tests were performed to determine whether distributions were significantly different from zero as these do not assume data are normally distributed. Variability was similar between the groups that were statistically compared. The Bonferroni method was used to correct for multiple comparisons. No statistical method was used to estimate sample size, but it was consistent with previous publications.

Molecular and biochemical results are presented as mean + s.e.m. Percentages stated are group means. All statistical analysis used Prism GraphPad software. Normality was determined using the D'Agostino and Pearson omnibus normality test. Variability was similar between the groups that were statistically compared. Comparison data for normally distributed data consisting of two groups were analysed by two-tailed unpaired *t*-tests. Comparison of normally distributed data consisting of three or more groups was by one-way ANOVA followed by Tukey's multiple comparisons test. Comparisons for non-normally distributed data were performed using Mann–Whitney tests. The statistical test, exact *P* values, and sample size (*n*) for each experiment are specified in the figure legend. For optogenetic ELISA data, two-sided unpaired Student's *t*-tests were performed to compare mice from the same litter that received different conditions. No statistical method was used to estimate sample size, but it is consistent with previous publications. Molecular and biochemical analysis used a minimum of three biological replicates per condition.

Data availability. Data are publicly available upon request from the corresponding author.

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ARTICLE RESEARCH



Extended Data Figure 1 | See next page for caption.



Extended Data Figure 1 | 5XFAD mice have reduced power in gamma during hippocampal SWRs. a, Mouse in virtual reality environment. b, LFP recorded in CA1, above, filtered for theta (left) or SWRs (right), middle, and gamma, below. c, Mean and standard deviation of the normalized power spectrum during theta. Each animal's power spectral density was normalized to its peak (n = 6 mice per group). **d**, Normalized power spectral densities during theta periods in 3-month-old 5XFAD (green, n = 6 mice) and WT (black/grey, n = 6 mice) mice. Each animal's power spectral density was normalized to its peak (in theta). e. Average SWR-triggered spectrograms for one WT and one 5XFAD mouse shows an increase in the gamma band during SWRs. This increase is lower in the 5XFAD mouse than in the WT mouse (n = 370 and 514 SWRs in WT and 5XFAD, respectively; WT mouse shown here is the same as in Fig. 1a). This range of frequencies is often called 'slow gamma' to distinguish it from faster oscillations (65-140 Hz) that have also been included in the gamma range but for which the origins are less well understood. f, Distributions for each recording (left) and the mean and standard error across sessions (right) of instantaneous gamma frequencies during SWRs in 5XFAD (green) and WT (black) mice show distributions around 40 Hz (n = 820, 800, 679, 38, 1,875, 57 gamma cycles per session in six 5XFAD animals in six recording sessions and 181, 1,075, 919, 1,622, 51, 1,860, 1,903 gamma cycles per session in six WT animals in seven recording sessions). g, Cumulative distribution of the z-scored gamma power during the 100 ms around the peak of the SWR for WT (black) and 5XFAD animals (green) for each animal (left) and the mean and standard error (shaded) across animals (right) (n = 514, 358, 430, 22, 805, 37 SWRs per session in six 5XFAD animals and 82, 311, 370, 776, 18, 710, 818 SWRs per session in six WT animals). h, Fraction of spikes in CA1 during SWRs as a function of the phase of gamma in 5XFAD (green) and WT (black) mice for each animal (left) and the mean and standard error across animals (right, *n* = 2475, 1060, 3092, 25, 6521, 123 spikes during SWRs per session in six 5XFAD mice and 360, 4741, 1,564, 2,961, 88, 3,058, 4,270 spikes during SWRs per session in six WT mice). i, SWR rate per non-theta period in 5XFAD (green) and WT (black) mice for each animal (left) and all animals combined (right, rank-sum test, $P < 10^{-10}$, n = 117, 210, 151, 55, 100, 1 non-theta periods per session in six 5XFAD mice and 80, 68, 115, 95, 15, 159, 218 non-theta periods per session in six WT mice). j, The cumulative distribution of gamma power during large SWRs (detection threshold greater than six standard deviations above the mean, Methods) shows significantly smaller increases in 5XFAD (green) than WT (black) mice (rank-sum test, $P < 10^{-5}$, n = 1,000 SWRs in six 5XFAD mice and 1,467 SWRs in 6 WT mice). k, Fraction of spikes as a function of the phase of gamma during large SWRs (detection threshold greater than six standard deviations above the mean, Methods), mean \pm s.e.m. (left) and histogram of the depth of modulation of spiking (right) as a function of gamma phase in 3-month-old 5XFAD (green, n = 6 mice) and WT (black, n = 6 mice) mice (rank-sum test, bootstrap resampling

 $P < 10^{-10}$, n = 2500 5XFAD spike-gamma phase distributions and 3,000 WT distributions). I, Power spectral density during 40 Hz stimulation (red, left), random stimulation (blue, centre), or no stimulation (black, right) of FS-PV-interneurons in CA1 for each mouse (n = 4 5XFAD mice with 169, 130, 240, 73 40 Hz, 143, 129, 150, 72 random, and 278, 380, 52, 215 no stimulation periods per animal and n = 3 WT mice with 65, 93, 91 40 Hz, 64, 93, 90 random, and 187, 276, 270 no stimulation periods per animal). m, Above: example raw LFP trace (above) and the trace filtered for spikes (300-6,000 Hz, below), with spikes indicated with red stars after optogenetic stimulation (blue vertical lines). Below: histogram of spikes per pulse after the onset of the 1 ms laser pulse during 40 Hz stimulation (red), random stimulation (blue), or no stimulation (black, n = 345,76240 Hz, 301559 random pulses, and 32,350 randomly selected no stimulation times at least 500 ms apart from 552 40 Hz, 543 random, and 1681 no stimulation periods in four 5XFAD and three WT mice). n, Histogram of the difference in firing rates between 40 Hz stimulation and random stimulation periods shows that both types of stimulation elicit similar amounts of spiking activity (Wilcoxon signed rank test for zero median, P > 0.6, n = 538 stimulation periods from four 5XFAD and three WT mice, NS, not significant). o, Multiunit firing rates per 40 Hz stimulation (red), random stimulation (blue), and no stimulation (black) period for each animal. Box-and-whisker plots show median (white lines in box) and quartiles (top and bottom of box). In all animals firing rates between 40 Hz and random stimulation were not significantly different, showing that the random stimulation condition serves as a control for spiking activity (rank-sum tests for each animal, three WT and four 5XFAD mice, P > 0.09, n = 87, 130, 8, 65, 93, 91, 73 40 Hz stimulation periods and 85, 129, 5, 64, 93, 90, 72 random stimulation periods per animal). We also examined whether 40 Hz stimulation caused neuronal hyperactivity relative to no stimulation, because, according to a recent report, this could have negative effects on neural circuit function²⁶. In most animals the firing rates between 40 Hz or random stimulation and no stimulation were not significantly different (rank-sum tests for each animal, two WT and two 5XFAD, *P* > 0.25, *n* = 8, 93, 91, 73 40 Hz stimulation periods and 15, 277, 270, 215 baseline periods per animal) or the firing rates during 40 Hz or random stimulation were lower than during no stimulation (rank-sum tests for each animal, one WT and one 5XFAD, $P < 10^{-5}$, which is significant when corrected for performing multiple comparisons, n = 130, 6540 Hz stimulation periods and 379, 187 baseline periods per animal) indicating that 40 Hz stimulation did not cause neuronal hyperactivity. In one animal there was significantly more activity with 40 Hz or random stimulation than during baseline (rank-sum test for one 5XFAD, mouse, $P < 10^{-5}$, n = 8740 Hz stimulation periods and 251 baseline periods per animal). Therefore in six out of seven animals we see no evidence that the 40 Hz optogenetic stimulation of FS-PV-interneurons causes hyperactivity.





ChR2-EYFP Hoechst



Extended Data Figure 2 | See next page for caption.

Rab5

Hoechst

Merge



Extended Data Figure 2 | ChR2 was expressed in FS-PV-interneurons for optogenetic stimulation. a, AAV-DIO-ChR2–eYFP or AAV-DIO–eYFP drives Cre-dependent expression of ChR2–eYFP or eYFP to produce cell-type-specific targeting of ChR2 or eYFP, respectively. In the presence of Cre, ChR2–eYFP or eYFP is inverted into the sense direction and expressed from the EF-1α promoter in PV-positive cells. ITR, inverted terminal repeat; polyA; WPRE, woodchuck hepatitis B virus post-transcriptional element. b, ChR2–eYFP was strongly expressed in PV-positive interneurons in CA1 of 3-month-old 5XFAD/PV-Cre mice (scale bar, 100 μm). c, Immunohistochemistry with anti-eYFP and anti-PV antibodies in CA1 of 3-month-old 5XFAD/PV-Cre mice expressing AAV-DIO-ChR2–eYFP shows eYFP expression only in PV-positive cells (scale bar, 50 μm). d, Representative western blots showing levels of full-length APP (top left, CT695), APP CTFs (bottom left, CT695), APP NTF (top right, A8967), and β -actin (bottom right, A5316, loading control) in CA1 in eYFP, random, and 40 Hz stimulation conditions, one mouse per lane, with two biological replicates of each condition. **e**, Immunohistochemistry with anti-A β (12F4, red) antibodies in CA1 of 5XFAD/PV-Cre mice expressing only eYFP or ChR2 with 40 Hz, and random stimulation conditions (scale bar, 50 µm). **f**, Bar graphs represent the relative immunoreactivity of A β normalized to eYFP (n = 4 mice per group; *P < 0.05, ***P < 0.001 by one-way ANOVA). Bar graphs show mean + s.e.m. **g**, Immunohistochemistry with anti-Rab5 antibody (ADI-KAP-GP006-E, green) in CA1 of 5XFAD/PV-Cre mice (scale bar, 50 µm). **h**, Relative Rab5 intensity levels normalized to eYFP controls (n = 3 mice per group).

a Up-regulated genes

2010002N04Rik	Beta-s	Card 10	Cyp2d22	Gide	H1fx	lqfbp4	Lag3	Mpped1	Pcsk1n	Prrt1	Sic12a9	Tmem198
2010300C02Rik	Bst2	Cd68	Dcakd	Gm129	H2-D1	Igfbpl1	Lcat	мп	Pdzd2	Rab40b	SIc25a34	Tpst2
2410018L13Rik	Clga	Cebpb	Egr4	Gm2115	H2-K1	Irt7	Lefty1	MI2	Pgls	Rara	Slc29a4	Trim30a
Adra2c	C1qb	Cebpd	Erf	Gng7	Hipk4	irf9	Lgals3b	Mtap1s	Phyhd1	Rasl11b	Spp1	Ttr
Agfg2	Ciqc	Cirbp	F730043M19Rik	Gpnmb	Hmox1	Itoka	Lingo3	Npy	Pitpnm2	Rbm3	Spsb1	Unc5a
Agxt211	C1ql2	Cnn2	Fam107a	Gpr25	1830012016Rik	Junb	Lrg1	Nr1d1	Plekhq5	Rpph1	Ssbp4	Ugcr11
Arc	C1qtnf4	Cotl1	Fam163b	Gpr3711	lcam1	Kcnc4	Ltbp4	Nr4a1	Pnpla7	Rprml	Sstr4	Usp18
Atf3	C3ar1	Crip2	Fmo2	Grm2	Icam5	Kcnh3	Lyz2	Oasl2	Pou3f1	Sbk1	Tfcp2l1	Vwf
B2m	C4b	Cst3	Fn1	Gstm1	lfit1	Kcnj4	Metrn	Palm	Ppp1r1a	Scara3	Thbs4	Wfs1
BC018242	Car7	Ctxn1	Gbp3	Gstm6	lfit3	KIF16	Mmp12	Parp14	Prr7	Sh3bgrl3	Thrsp	Xdh

b Down-regulated genes

1700003M02Rik	A630089N07Rik	Arhgap24	Codc141	Cited2	Edil3	Fam19a4	Gpr115	Hs6st2	Kenma1	Lrrc4B	Necabl	Pcsk1	Prkcd	Rgs16	Sh3bgrl2	Spata18	Th	Tyrp1	Ysk4
1700007K13Rik	AF529169	Asb2	Codc153	Clec18a	Efcab1	Fbln7	Gpr123	Hsp90aa1	Kann3	Lirc55	Necab3	Pdp1	Prkch	Rgs3	Shox2	Spock1	Timp2	Ubxn10	Zechc12
1700009P17Rik	AU021034	Aspa	Code19	Clic6	Efna5	Faf1	Gpr139	Hsp90b1	Kong10t1	Malat1	Nexn	Peg10	Prkcg	Rgs4	Shroom3	Spock3	Tm4sf1	Ugt8a	Zdb/Z
1700026D08Rik	AW551984	Baiap3	Code3	Cntn6	Eif5a2	Fqf10	Gpr151	Hspa4I	Kctd12b	Mcf2	Nfam1	Pgap1	Prkq2	Rgs6	SIc12a2	Srgap1	Tmem130	Une13c	Zdhhc22
1700027A23Rik	Adamts 15	Bbox1	Code40	Cntnap4	Elavi2	Fhdct	Gpr153	Hir2c	Kctd8	Megf11	Ngb	Pgbd1	Prokr2	Rims3	SIc17a6	St8sia2	Tmem132c	Vang[1	Zfhx3
1700028P14Rik	Adamts9	Bmp7	Codic81	Cobl	Elavi4	Eoug1	Gpr26	Htr5b	Kif9	Mgat4c	Nhlh2	Phactr2	Pn5	Rit2	Stc38a1	Strbp	Tmem163	Vat1	Zfp474
1700040L02Rik	Adcyap1	Blbd11	Cd109	Coch	Elfn1	Foxp2	Gpr4	Htr7	Kit	MIT	Nppa	Pirt	Prrg4	Rnf152	SIc39a4	Sv2b	Tmem176a	Vat1I	Zfp618
1700094D03Rik	Adra1b	C530008M17Rik	Cd24a	Col12a1	Emb	Frem3	Gprasp2	Hydin	Kitt.	Mme	Npr1	Pkib	Ptods	Robo1	SIc5a3	Sympo2	Tmem212	Vav2	Zfp941
1810041L15Rik	Aebp1	Cacha2d2	Cdh26	Col8aZ	Enkur	Fstl5	Gpx3	Inadl	Kihi1	Mob3b	Nr4a2	Plag11	Pton14	Rorb	SIc5a7	Syt15	Tmem56	Vav3	Zic1
2310039L15Rik	Agt	Calb2	Cdhra	Cpne4	Eno4	Fzd1	Grb10	lgea	Lars2	Mreq	Nrip3	Plcb4	Pvrl2	Rps6ka6	SIc6a9	Syt4	Tric	Wwa5b1	Zic2
2410004P03Rik	Alf1	Calr3	Cdr1	Cone9	Enox2	Fzd10	Gria4	gub	Lbh	Mrvi1	Nrp2	Pich1	Pvrl3	Rsph1	Sic7a11	Syt6	Tont1	Wbscr27	Zic3
3110047P20Rik	Ak4	Camk2d	Cdr2	Dach1	Epha8	Galn(13	Grid2ip	lox1	Lbp	Msi2	Nrsn2	Plch2	Rab37	Rsph4a	Sic9a4	Syt9	Trhde	Wdr16	Zic4
3632451006Rik	Ak7	Car10	Chat	Don	Eph3	Gap43	Gnn3a	Irx2	Ldhd	Mtfr1	Ningt	Plcxd2	Ramp3	Rspo2	Slco2a1	Tac1	Trim36	Wdr52	Zic5
4930451C15Rik	Akap12	Cast	Chab	Ddc	Ermn	Gatm	Grk4	Irx3	Lect1	Mumili	Nudt4	Pld5	Ranbp3/	Scnta	Slit2	Tac2	Trim66	Wdr6	1
4932411L15	Amigo2	CbIn1	Chrdl1	Dnahc5	Etv1	Gdpd5	Grm4	Itga3	Lef1	Musk	Olfm3	Plekha1	Rapigap	Scube1	Siltrk6	Tacr1	Trpc3	Wdr78	
4932425124Rik	Amoti 1	Cbln2	Chrna3	Dnahc6	Exph5	Gfra1	Gucy1a:	Kcnc2	Lhfpl	Myb	Optn	Planc1	Rasgef1b	Scube3	Sncg	Tcf712	Trps1	Wdr96	
5730508B09Rik	Ankrd29	Cbin4	Chma4	Dpp10	Fabp7	Gm6300	Hcn4	Kong4	Lhfpl3	Mycbpap	Ob/2	Popdc3	Rass19	Sema3d	Sntn	Tekt1	Tsku	Wfikkn2	1
6330406115Rik	Ankrd34c	Ccdc108	Chmb3	Dpy19l1	Fam149a	Gm7609	Hdc	Konip1	Lhx9	Мурс	Pamri	Pou4f1	Rbms3	Sema6a	Socs2	Tex15	Ttc18	Wift	
A2m	Ano1	Ccdc135	Chrnb4	Dynlrb2	Fam196b	Gm973	Hhip	Kcni12	Lrquk.	Ndn	Pbx3	Ppp1r32	Resp18	Serpinf1	Sox5	Tex9	Ttc21a	Wis	·
A330021E22Rik	Aup4	Ccdc136	Cit	Ebf1	Fam198b	Gng8	Hivep1	Koni16	Linc23	Necab1	Pcp4	Ppp1r36	Ret	Sopp2	Spag16	Tafb2	Ttc39a	Wnt3	11 11 1





d

Enrichment of up-regulated genes under different perturbations

Name of perturbed transcriptome	NES	Norminal p-value	FDR q-value
MCSF treated microglia	1.76	0.000	0.000
NMDA treated neurons	1.62	0.000	0.000
IL34 treated microglia	1.59	0.000	0.000
GMCSF treated microglia	1.49	0.005	0.004
Bicuculline treated neurons	1.49	0.016	0.013
ALS SOD1 mutant microglia	-1.26	0.050	0.028
LPS&IFNg treated macrophage (M1)	1.18	0.122	0.081
MeCP2 null microglia	1.16	0.164	0.127
IL4 treated macrophage (M2)	-1.19	0.101	0.147
Huntington HTT mutant microglia	1.03	0.371	0.361
Germ-free microglia	0.94	0.604	0.667
Talendologin Instead nourons	0.76	0.941	0.970

NES: normalized enrichment score







Extended Data Figure 3 | See next page for caption.



Extended Data Figure 3 | Optogenetically driven 40 Hz oscillations in CA1 cause changes in gene regulation and immediate early gene expression. a, Table of 130 genes upregulated by 40 Hz FS-PV-interneuron stimulation determined by whole-transcriptome RNA-seq of CA1 from 3-month-old 5XFAD/PV-Cre mice (P < 0.05 by Cufflinks 2.2). b, Table of 393 genes downregulated by 40 Hz FS-PV-interneuron stimulation determined by whole transcriptome RNA-seq of CA1 from 3-month-old 5XFAD/PV-Cre mice (P < 0.05 by Cufflinks 2.2). c, Box plot showing FPKM values of up- and downregulated genes in eYFP and 40 Hz groups. Box shows median (black lines in box) and quartiles (top and bottom of box), whiskers show minimum and maximum values, and circles show outliers. d, GSEA statistics tables showing statistical significance of correlation between genes up- or downregulated by 40 Hz stimulation and publicly available neuron-, microglia-, and macrophage-specific RNA-seq data under different chemical and genetic perturbations; the perturbation terms were ranked on the basis of the false discovery rate *q* values for the upregulated gene list, from the smallest to the largest (Methods). **e**, RT-qPCR verification of specific gene targets in the RNA-seq dataset. Bar graph shows relative RNA levels (fold change) from eYFP (black) and 40 Hz stimulation (red) conditions (*P < 0.05, **P < 0.01, ***P < 0.001 by Student's *t*-test, n = 3 mice per group). All bar graphs show mean + s.e.m. **f**, Immunohistochemistry with Hoechst to label all cell nuclei in CA1 of 5XFAD/PV-Cre mice expressing only eYFP or ChR2 with 40 Hz stimulation conditions (scale bar, 50 µm). **g**, Bar graph represents the estimated CA1 thickness in 5XFAD/PV-Cre mice expressing only eYFP or ChR2 with 40 Hz stimulation conditions (n = 4 mice per group; NS, not significant by Student's *t*-test).



Extended Data Figure 4 | A 40 Hz light flicker drives 40 Hz oscillations in VC, while random flickering does not. a, Power spectral densities of LFPs in VC during 40 Hz light flicker (red, far left), random light flicker (blue, centre left), dark (black, centre right), or light (green, far right) in VC for each recording session for each mouse (n = 5 recordings from four 5XFAD mice with 47, 51, 64, 49, 16 40 Hz flicker, 47, 50, 64, 50, 16 random flicker, 279, 301, 382, 294, 93 dark and 47, 50, 64, 49, 15 light periods). Light flicker at other frequencies increased power in the flicker frequency, as others have found previously^{23,24} (data not shown). **b**, Histogram of the difference in firing rates between 40 Hz light flicker and random light flicker (n = 226 stimulation periods from five recording sessions in four 5XFAD mice). c, Multiunit firing rates in VC during 40 Hz light flicker (red), random light flicker (blue), dark (black), or light (green) periods. Box plots show median (white lines in box) and guartiles (top and bottom of box). In all animals, firing rates between 40 Hz flicker and random flicker conditions were not significantly different showing that the random stimulation condition serves as a control for spiking activity (rank-sum tests for each of five recording session from four 5XFAD mice, P > 0.06, n = 47, 51, 64, 49, 16 40 Hz flicker periods and 47, 50, 64, 50, 16 random flicker periods per recording). There were no significant differences in firing rates between 40 Hz flicker and light conditions, indicating that

40 Hz light flicker generally did not cause neuronal hyperexcitability (rank-sum tests for each of five recording sessions from four 5XFAD mice, P > 0.2 for four recording sessions, P < 0.01 for one recording session, which is not significant when corrected for performing multiple comparisons, *n* = 47, 51, 64, 49, 16 40 Hz periods and 47, 50, 64, 49, 16 light periods per recording). In one session, there was more activity in the 40 Hz flicker than in the dark condition. d, Example traces of LFPs recorded above the brain during light flicker (above, yellow indicates light on, black indicates light off), during three different recording sessions. e, Power spectral densities of LFPs recorded above the brain during 40 Hz light flicker show no increase in power at 40 Hz. Thus, the effect is not due to photoelectric effects on recording equipment or electrical noise (n = 4, 2, 1, 1, 17, 42, 36, 55, 53 40 Hz flicker periods from four recording sessions in three 5XFAD mice undergoing VC recordings and from five recording sessions in two 5XFAD and three WT mice undergoing hippocampal recordings). Mean (solid line) and standard deviation (shaded area) across recordings are shown on the left and per animal on the right. Recordings with fewer than three flicker periods (light red) resulted in noisier power spectral densities than recordings with more data (dark red) but none showed evidence of peaks at 40 Hz.

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Extended Data Figure 5 | A 40 Hz light flicker does not affect A blevels in hippocampus or barrel cortex. a, Example LFP trace in hippocampal CA1 before and during 40 Hz light flicker (above). Mean (solid line) and standard deviation (shaded area) of power spectral density during 40 Hz light flicker (red), random light flicker (blue), or dark (black) in CA1 (n = 25XFAD and 3 WT mice). **b**, Histogram of the fraction of spikes in hippocampus as a function of time for four cycles of 40 Hz light flicker (left, red) or the equivalent period of time for random light flicker (right, blue, n = 2 5XFAD and 3 WT mice, mean \pm s.e.m. across animals). Bar above indicates when light was on (yellow) or off (black). For random stimulation, spiking was aligned to the start of the light turning on; additional periods with light-on occurred at random intervals are indicated by grey (Methods). c, Histogram of the difference in firing rates between 40 Hz light flicker and random light flicker (bottom n = 168stimulation periods from five recording sessions in two 5XFAD and three WT mice). d, Power spectral densities of LFPs in CA1 during 40 Hz light flicker (red, far left), random light flicker (blue, centre left), dark (black, centre right), or light (green, far right) for each recording session for each mouse (n = 5 recordings from two 5XFAD and three WT mice with 22, 54, 42, 71, 55, 40 Hz flicker, 12, 34, 32, 54, 36 random flicker, 115, 240, 224, 342, 282 dark and 12, 33, 33, 54, 35 light periods). e, Multiunit firing rates in CA1 during 40 Hz light flicker (red), random light flicker (blue), dark (black), or light (green) periods. Box plots show median (white lines

in box) and quartiles (top and bottom of box). In all animals firing rates between 40 Hz flicker and random flicker conditions were not significantly different, showing that the random stimulation condition serves as a control for spiking activity (rank-sum tests for each of five recordings from two 5XFAD and three WT animals, P > 0.2, n = 22, 54, 42, 71, 5540 Hz flicker periods and 12, 34, 32, 54, 36 random flicker periods per recording). There were no significant differences in firing rates between 40 Hz flicker and light conditions indicating that 40 Hz light flicker generally did not cause neuronal hyperexcitability (rank-sum tests for each of five recordings from two 5XFAD and three WT animals, P > 0.3, n = 22, 54, 42, 71, 55 40 Hz periods and 12, 34, 33, 54, 35 light periods per recording). f, Bar graphs of relative $A\beta_{1-40}$ levels in VC of 5XFAD mice in dark, 40 Hz flicker, and random flicker conditions, normalized to dark (n = 4 mice per group; NS, not significant). Bars represent mean + s.e.m. Circles superimposed on bars in bar graphs indicate individual data points in each group. \hat{g} , Bar graphs of relative $A\beta_{1-42}$ levels in VC of 5XFAD mice in dark, 40 Hz flicker, and random flicker conditions, normalized to dark (n = 4 mice per group; NS indicates not significant). Bars represent mean + s.e.m. Circles superimposed on bars indicate individual data points in each group. **h**, Bar graph of relative $A\beta_{1-40}$ and $A\beta_{1-42}$ levels in barrel cortex of 5XFAD mice in dark and 40 Hz flicker conditions, normalized to dark (n = 3 mice per group; NS, not significant by Student's *t*-test).



Extended Data Figure 6 | Acute reduction in A β after light flicker in APP/PS1 and WT mice. a, Bar graph of relative A β_{1-40} and A β_{1-42} levels of APP/PS1 in VC in dark and 40 Hz flicker conditions, normalized to dark (n = 4 mice per group for dark and n = 4 mice per group for 40 Hz flicker conditions; NS, not significant, *P < 0.05, by Student's *t*-test). All bar graphs show mean + s.e.m. throughout this figure. Circles superimposed

on bars in bar graphs indicate individual data points in each group. **b**, Bar graph of relative mouse $A\beta_{1-40}$ and $A\beta_{1-42}$ levels in VC of 9-month-old WT mice in dark and 40 Hz flicker conditions, normalized to dark (n = 11 mice per group for dark and n = 9 mice per group for 40 Hz flicker conditions; *P < 0.05, by Student's *t*-test).

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Extended Data Figure 7 | A 40 Hz light flicker does not decrease synaptic density in VC. a, Schematic depicting isolation of microglia from VC. VC was dissected, then single cells were suspended and labelled with CD11b and CD45 antibodies. Subsequently, cells were sorted by FACS and lysed. A β_{1-40} levels were analysed by ELISA. b, Bar graph of A β_{1-40} levels in microglia purified using FACS (Methods) from VC of 3-month-old 5XFAD and WT mice (n = 8 mice per group for 5XFAD and n = 4 mice per group for WT mice; *P < 0.05 by Student's *t*-test).

Circles superimposed on bars indicate individual data points in each group. **c**, Immunohistochemistry with SVP38 (red) antibodies to detect synaptophysin in VC of 3-month-old 5XFAD mice in dark and 40 Hz flicker conditions (images were taken with 40× objective; scale bar, 50 μ m). Right: 100× rendering of dark and 40 Hz flicker conditions. **d**, Bar graph of relative SVP38 intensity levels in VC of 5XFAD mice after dark (black) and 40 Hz (red) flicker conditions, normalized to dark (n = 4 mice per group; NS, not significant by Student's *t*-test).



a Recovery after 1 hr 40 Hz flicker

Extended Data Figure 8 | A β levels in VC return to baseline 24 h after 1 h of 40 Hz light flicker. a, Bar graph of relative A β_{1-40} and A β_{1-42} levels in VC of 5XFAD mice 1, 4, 12, and 24 h after 1 h of dark or 40 Hz flicker treatment, normalized to dark (n = 4 mice per group for 4 and 12 h wait, n = 6 for 1 and 24 h wait, n = 12 for dark; NS, not significant, *P < 0.05, **P < 0.01 by one-way ANOVA).



Iba1

Hoechst

Extended Data Figure 9 | Driving 40 Hz oscillations in VC via light flicker reduces phosphorylated tau in a tauopathy mouse model. a, Immunohistochemistry with anti-pTau (S202, green) and anti-MAP2 (red) antibodies in VC of 4-month-old P301S mice after 7 days of 1 h per day dark or 40 Hz flicker conditions (images were taken with $40 \times$ objective; scale bar, 50 μ m). Insets: 100 \times rendering of representative cell body in dark and 40 Hz flicker conditions. No changes were observed by western blot (data not shown). b, Bar graph of relative pTau (S202) intensity levels in P301S mice after 7 days of 1 h per day dark (black) and 40 Hz flicker (red) conditions (n = 8 mice per group; *P < 0.05 by Student's *t*-test). All bar graphs show mean + s.e.m. throughout. **c**, Bar graph of relative MAP2 intensity levels in P301S mice after 7 days of 1 h per day dark (black) and 40 Hz flicker (red) conditions (n = 8 mice per group; NS, not significant by Student's t-test). d, Immunohistochemistry with antipTau (S396, red) antibodies in P301S mice after 7 days of 1 h per day dark and 40 Hz flicker conditions (scale bar, 50 µm). e, Bar graph of relative pTau (S396) fluorescence intensity levels in P301S mice after 7 days of 1 h per day dark (black) and 40 Hz flicker (red) conditions (n = 8 mice per group; ****P < 0.0001 by Student's *t*-test). **f**, Immunohistochemistry with

anti-pTau (S404, green) antibodies in P301S mice after 7 days of 1 h per day dark and 40 Hz flicker conditions (scale bar, 50 µm). g, Bar graph of relative pTau (S400/T403/S404) fluorescence intensity levels in P301S mice after 7 days of 1 h per day dark (black) and 40 Hz flicker (red) conditions (n = 8 mice per group; **P < 0.01 by Student's t-test). Bar graphs show mean + s.e.m. h, Immunohistochemistry with anti-Iba1 (019-19741, green) antibodies in 4-month-old P301S mice after 7 days of 1 h per day dark and 40 Hz flicker conditions (images were taken with 40× objective; scale bar, 50 μ m.) Insets: 100 \times rendering of representative microglia in eYFP and 40 Hz stimulation conditions. i, Bar graph of the number of microglia in P301S mice after 7 days of 1 h per day dark (black) and 40 Hz flicker (red) conditions (n = 8 mice per group; NS, not significant by Student's t-test). j, Bar graph of the diameter of microglia cell bodies in P301S mice normalized to dark after 7 days of 1 h per day dark (black) and 40 Hz flicker (red) conditions (n = 8 mice per group; ****P < 0.0001 by Student's t-test). k, Bar graph of the average length of microglia primary processes in P301S mice normalized to control after 7 days of 1 h per day dark (black) and 40 Hz flicker (red) conditions (n = 8 mice per group; *****P* < 0.0001 by Student's *t*-test).

Extended Data Table 1 | Raw A $\!\beta_{1\text{--}40}$ and A $\!\beta_{1\text{--}42}$ concentrations

Treatment Dilution Factor		Average Aβ ₁₋₄₀ Concentration (pg/ml)	Average $A\beta_{1,42}$ Concentration (pg/ml)			
Optogenetics		1				
PV-Cre EYFP	1:2	100.01, 61.598, 65.462, 82.509, 69.023, 70.831, 82.152, 74.314	58.777, 54.546, 30.585			
PV-Cre 40 Hz	1:2	46.604, 31.041, 26.639, 55.612, 69.326, 17.711, 3.9951	27.271.41.950.18.790,18.262			
PV-Gre 8 Hz	1:2	101.268, 54.283, 90.190, 151,690	50,699, 122,85, 35,507			
PV-Cre Random	1:2	235.68, 89.962, 157.37, 323.902, 451.78, 241.63	54,029, 137,78, 144.63			
oCaMKII-Cre EYFP	1:2	45.813, 59.069, 40.404, 66.810	72.052, 36.573, 67.243, 59.295			
oCaMKII-Cre 40 Hz:	1:2	55.942. 44.270, 57.498, 47.382, 115.08, 75.673	70.847, 79.683, 61.429			
aCaMKII-Cre 8 Hz	1:2	52.829, 46.604, 57.720	95.939, 21.640, 102.987			
cCaMKII-Cre Random	1:2	218.00, 191.72, 159.07	66.203, 168.867, 176.404			
Light flicker						
Dark 1 hr VC	1;2	343.8, 245.3, 210.6, 343.8, 588.4, 394.9, 123.3, 336.3, 328.2, 579.1, 420.0, 339.2	449.5, 320.7, 275.2, 449.5, 769.2, 516.2, 449.4, 320.6, 275.2, 449.4, 769.1, 516.1			
Light 1 hr VC	1:2	366.9, 632.4, 378.2, 314.1, 266.9, 264.1	616.4, 592.3, 802.9, 394.5, 330.7, 337.8			
20 Hz 1 hr VC	1:2	944.4, 313.2, 595.9, 530.0, 456.5, 289.9	1624, 302.4, 816.0, 687.2, 676.6, 343.0			
40 Hz 1 hr VC	1:2	146.4, 143.6, 104.9, 99.6, 179.7, 219.8	191.4, 187.7, 137.2, 130.2, 234.9, 287.3			
80 Hz 1 hr VC	1:2	332.5, 328.7, 363.5, 390.6, 530.0, 673.3	558.3, 418.9, 510.7, 609.5, 1186, 921.9			
40 Hz + PTX 1 hr VC	1:2	367.2, 431.4, 445.2, 392.4, 386.7, 445.2	396.6, 540.5, 532.7, 705.0, 104.5, 104.5			
Random 1 hr VC	1:2	461.8, 100.2, 9.819, 416.6	423.9, 157.9, 389.9, 841.5			
Dark 1 hr HPC	1:2	97.949, 107.33, 119.92, 139.33	499.30, 355.13, 469.53, 598.03			
40 Hz 1 hr HPC	1:2	88.136, 104.78, 161.52, 197.36	364.53, 408.41, 436.62, 873.83			
Random 1 hr HPC	1:2	95.816, 136.77, 70.004, 125.47	466.39, 500.87, 311.26, 582.355			
Dark 7 days soluble	1:50	1216.9, 1181.3, 1173.4, 1199.5, 134.73, 151.34, 113.26, 145.14, 127.91, 127.48, 143.02, 127.48, 141.07	5217.2, 8057.9, 9051.3, 6773.7, 244.11, 236.96, 235.38, 240.62, 286.19, 8.382, 11.21, 14.03, 13.56			
Dark 7 days insoluble	1:100	1173.2, 1208.2, 1205.3, 1214.6, 994.86, 1059.2, 1176.6, 1065.4, 1002.9, 306.16, 690.70, 3442.7, 152.73	8572.7, 9127.1, 6349.3, 10138, 6852.2, 7056.7, 7039.7, 7094.2, 7289.0, 748.21, 1117.1, 1055.5, 504.95			
40 Hz 7 days soluble	1:50	476.71, 283.83, 336.87, 237,22, 7.0175, 4.1480, 4.0580, 1.5205, 91.864, 152.73, 148.84, 141.07, 162.44	419.7, 248.1, 242.7, 90.974, 95.626, 56.936, 67.577, 47.586, 200.87, 13.56, 9.794, 15.44, 3.677			
40 Hz 7 days insoluble	1:100	281.97, 270.37, 86.199, 239.71, 23.557, 15.166, 22.714, 1038.9, 1099.8, 1760.8, 1558.8, 187.69, 22.64	202.96, 130.71, 195.73, 193.70, 1646.89, 1579.1, 503.44, 1400.0, 7536.62, 955.23, 1208.8, 694.57, 784.91			
Dark 1 hr BC	1:2	81.874, 18.343, 86,554	391.95, 883,69, 604.97			
40 Hz 1 hr BC	1:2	81.307, 27.986, 30.113	300.34, 1152.5, 616.92			
Dark 1 hr	1:2	151.4, 334.4, 301.1, 185.6	223.3, 423.2, 375.0, 250.9			
40 Hz 1 hr wait 1 hr	1:2	97.6, 95.8, 70.0, 66.4, 119.9, 146.6	43.3, 44.8, 35.45, 35.35, 70.5, 90,4			
40 Hz 1 hr wait 4 hr	1:2	91.06, 141.8, 111.2, 12.30	108.0, 168.1, 157.3, 35.158			
40 Hz 1 hr wait 12 hr	1:2	167.2, 101.6, 89.31, 119.9	236.1, 134.6, 124.8, 152.4			
40 Hz 1 hr wait 24 hr	1:2	246.7, 177.6, 281.2, 175.0, 257.3, 204.2	231.8, 107.0, 402.7, 184.6, 245.1, 179.7			
Dark APP/PS1	1:2	1050.16, 1085.25, 1522.45, 1153.69, 1750.77	19.22, 30.68, 28.08, 14.25, 25.30			
40 Hz APP/PS1	1:2	512.42, 947.80, 850.45, 793.63	18.85, 15.58, 18.92, 11.44			
Dark WT	1:1	0.038, 0.813, 2.016, 1.913, 0.313, 4.11, 7.23, 20.2, 40.4, 38.7, 11.9	N/A			
40 Hz WT	61	0.139, 0.325, 0.346, 0.390, 8.92, 12.1, 6.34, 12.4, 13.1	N/A			

Raw A β_{1-40} and A β_{1-42} levels with ELISA dilution ratios for each experimental group. Equal tissue masses were compared for each ELISA experiment. For 7-day experiments, values were normalized to within litter controls such that raw values 1–4 in each condition were normalized to the mean of 'Dark' values 1–4; raw values 5–9 in each condition were normalized to the mean of 'Dark' values 5–9; raw values 10–13 in each condition were normalized to the mean of 'Dark' values 10–13.

CORRECTION

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Author Correction: Gamma frequency entrainment attenuates amyloid load and modifies microglia

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In Extended Data Fig. 8 of this Article, we inadvertently copied the data between A β_{1-40} levels at 1 h and A β_{1-42} levels at 1 h. We have corrected the A β_{1-42} graph and re-run the statistical analysis. We see a significant reduction in A β_{1-40} and A β_{1-42} levels 1 h after 40 Hz flicker, consistent with the other independent replications in the Article and with our immunohistochemistry analysis of A β plaques. Our overall findings and conclusions are not changed by these results. We also accidentally omitted the raw ELISA data values for A β_{1-40} and A β_{1-42} , for 'Dark 1hr' and '40 Hz 1 hr wait 1 hr' in Extended Data Table 1. The Supplementary Information to this Amendment contains the old, incorrect Extended Data Fig. 8 and Extended Data Table 1, for transparency. These errors have been corrected online.

Supplementary information is available for this Amendment at https://doi. org/10.1038/s41586-018-0351-4.