Early death in a mouse model of Alzheimer’s disease exacerbated by microglial loss of TAM receptor signaling

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Recurrent seizure is a common comorbidity in early-stage Alzheimer’s disease (AD) and may contribute to AD pathogenesis and cognitive decline. Similarly, many mouse models of Alzheimer’s disease that overproduce amyloid beta are prone to epileptiform seizures that may result in early sudden death. We studied one such model, designated APP/PS1, and found that mutation of the TAM receptor tyrosine kinase (RTK) Mer or its ligand Gas6 greatly exacerbated early death. Lethality was tied to violent seizures that appeared to initiate in the dentate gyrus (DG) of the hippocampus, where Mer plays an essential role in the microglial phagocytosis of both apoptotic and newborn cells normally generated during adult neurogenesis. We found that newborn DG neurons and excitatory synapses between the DG and the cornu ammonis field 3 (CA3) field of the hippocampus were increased in TAM-deficient mice, and that premature death and adult neurogenesis in these mice were coincident. In contrast, the incidence of lethal seizures and the deposition of dense-core amyloid plaques were strongly anticorrelated. Together, these results argue that TAM-mediated phagocytosis sculpts synaptic connectivity in the hippocampus, and that seizure-inducing amyloid beta polymers are present prior to the formation of dense-core plaques.

Alzheimer’s disease (AD), the most prevalent neurodegenerative disease and most common cause of human dementia, is associated with seizures. Unprovoked seizures are substantially more frequent in sporadic AD patients compared to age-matched nondemented populations (1–6) and are exceptionally prominent in early stages of early-onset familial AD (4, 7, 8). In addition, subclinical epileptiform activity has been detected by long-term electroencephalography (EEG) in more than 40% of AD patients, where it is correlated with accelerated cognitive decline (9). This abnormal network activity often begins decades before the symptomatic onset of AD (10, 11). Correspondingly, many amyloidogenic mouse models of AD that rely on the elevated production of amyloid beta (Aβ) peptides—as a result of transgenic expression of mutant forms of the human amyloid precursor protein (APP) and/or the presenilin 1 (PSEN1) component of the APP gamma secretase—also display cortical and hippocampal hyperexcitability and experience unprovoked seizures early in disease before the appearance of Aβ plaques (12–19).

We have recently published analyses of one such mouse—the APP/PS1 line that carries an amyloidogenic “Swedish” mutation in the human APP gene and a pathogenic mutation in the human PSEN1 gene (20–22). Although not routinely addressed in the literature, it is well documented that APP/PS1 populations on a C57BL/6 background exhibit 10 to 15% mortality due to sudden death early in life (12, 13). This sudden death, which is not associated with any gross postmortem neuropathology, has been found to be the result of lethal epileptic seizures (12). These seizures are in turn linked to markedly higher levels of interictal spiking of cortical pyramidal neurons, as detected by EEG, patch clamp recording in cortical slices, and implanted electrodes in vivo (12).

In our previous studies (22), we examined the effects of germline loss-of-function mutations in the mouse genes encoding the TAM receptor tyrosine kinases (RTKs) Mer and Axl (23, 24), and their ligand Gas6 (25), when crossed into APP/PS1. In the current analysis, we found that mutation of Mer (gene name MerTK), Gas6, or Axl and Mer together all led to a marked exacerbation of early sudden death in both sexes. Importantly, this was also seen for APP/PS1 compound mutants that carried microglial-restricted mutations in MerTK. As for APP/PS1 alone, nearly all sudden death in the compound mutants occurred before postnatal day (P)150, after which time all populations were stable. Video recording in our colonies demonstrated that the compound mutants died from the same spontaneous epileptic seizures that have been shown to kill young APP/PS1 mice. Analyses of the activation marker Arc immediately following lethal seizures were consistent with these seizures being initiated in the dentate gyrus (DG) of the hippocampus.

Significance

Most patients with Alzheimer’s disease (AD), a neurodegenerative disorder that leads to severe cognitive decline, suffer from comorbid seizures. These events also occur in mouse models of AD in which amyloid beta peptides are overproduced and are often lethal. Here we report the surprising finding that mutations in the TAM receptor tyrosine kinase system greatly exacerbate the incidence of lethal seizures in the APP/PS1 mouse AD model. This exacerbation is tied to TAM activity in microglia during adult neurogenesis in the hippocampus, a trigger site for seizures generally, and results in the development of exuberant excitatory synapses in this critical brain region.
Together with the subventricular zone (SVZ) adjacent to the lateral ventricle, the DG harbors a prominent site of adult neurogenesis in the mouse (26, 27). At both sites, microglial Mer has been shown to play an essential role in the phagocytic clearance of the large number of apoptotic cells (ACs) that are generated during this process (28, 29). Importantly, the bulk of adult neurogenesis and AC generation in the mouse occurs over a delimited developmental window between birth and P150 (26), the same window in which APP/PS1 mice die from seizures. During this time, Mer-expressing microglia also phagocytose live newborn cells that express the “eat-me” signal phosphatidylserine (PtdSer) (29, 30). In the SVZ, TAM mutations have been shown to result in the survival of many of these newborn cells, which would otherwise have been “eaten alive,” and in their integration as viable differentiated neurons in the olfactory bulb (29). Consistent with these earlier findings, we observed that APP/PS1 mice lacking both Axl and Mer displayed 6-fold more ACs and 2.1-fold more viable 5-bromo-2-deoxyuridine (BrdU)-labeled cells in the DG at 4 wk after a BrdU pulse than did APP/PS1 mice alone. Moreover, these TAM-deficient APP/PS1 mice also displayed a 1.4-fold increase, relative to their APP/PS1 counterparts, in excitatory connections between DG granule cell neurons and their synaptic targets in the cornu ammonis field 3 (CA3) of the hippocampus. Taken together, these results indicate that the loss of TAM signaling exacerbates the preexisting hyperexcitability of APP/PS1 mice via increased excitatory connections between the DG and CA3, and that TAM-mediated microglial phagocytosis is normally required for the proper sculpting of these hippocampal circuits during adult neurogenesis.

Results

Early Lethality in APP/PS1 Is Exacerbated by Merk and Gas6 Mutation. We monitored and plotted the survival of APP/PS1 mice on C57BL/6J over the first 200 d of postnatal life and quantified the loss of ~15% of the APP/PS1 colony by P150 (Fig. 1). This is in close agreement with previous reports of early sudden death in this line, which has been linked unequivocally to lethal tonic-clonic seizures (12, 13). For example, recordings from electrodes implanted into both APP/PS1 cortices have consistently detected profound and chronic epileptiform activity, with 65% of APP/PS1 mice experiencing at least one large electrographic seizure by 19 wk of age (12). Similarly, 24-h video monitoring has revealed that 83% of compound APP/PS1 mutants experiencing a grand mal seizure die immediately from the observed seizure (13). Treatment with the anticonvulsant levetiracetam, which suppresses seizures, entirely suppresses early sudden death in these mice; and conversely, treatment with the noncompetitive GABA\(_\mathrm{A}\) inhibitor picrotoxin, which antagonizes inhibitory neuronal signaling, increases seizure activity and exacerbates early sudden death (13). In contrast to APP/PS1 mice, mutants lacking both Mer and Axl (Axl~\(^{-}\)Merk~\(^{-}\)), the TAM receptors expressed by microglia (22, 29), together with mice lacking the universal TAM ligand Gas6 (Gas6~\(^{-}\)) (25, 31), all displayed normal wild-type (WT) (100%) viability well beyond P200 (Fig. 1), also as reported previously (23, 24, 32).

Although TAM system mutations on their own were without effect on viability, crossing these mutations into the APP/PS1 line greatly exacerbated its early sudden death. Introducing the Axl~\(^{-}\) single mutation yielded a trend toward increased lethality, but this effect was not statistically different from the course of death seen in APP/PS1 alone when evaluated using the log-rank test (Fig. 1; \(P = 0.3012\)). In marked contrast, introduction of Merk~\(^{-}\) or Axl~\(^{-}\)Merk~\(^{-}\) or Gas6~\(^{-}\) mutations into APP/PS1 all resulted in the death of ≥40% of the compound mutants by P150, and this exacerbation of early lethality relative to APP/PS1 was highly significant (Fig. 1). Sudden death in APP/PS1Merk~\(^{-}\), APP/PS1Axl~\(^{-}\)Merk~\(^{-}\), and APP/PS1Gas6~\(^{-}\) mice was first observed at or just before P50 and progressed at the same consistent rate in all three genotypes until ~P120 (Fig. 1). After ~P150 (dotted vertical line in Fig. 1), sudden death in all populations largely ceased. This has also been previously documented for APP/PS1 alone, whose populations are subsequently stable (100% viable) through at least 10 mo (13).

The two most prominent cellular loci for Mer expression in the adult brain are microglia, the specialized tissue macrophages of the central nervous system, and endothelial cells of the blood vasculature (22, 29, 33). To assess the relative importance of these two cell types to the APP/PS1Merk~\(^{-}\) lethality reported in Fig. 1, we crossed conditional floxed alleles of the Merk gene (29) into a tamoxifen-inducible Cx3cr1\(^{CreER}\) driver line (34) and then crossed these mice with APP/PS1. The Cx3cr1\(^{CreER}\)Merk~\(^{f/f}\) mutants allow for tamoxifen-induced microglial Merk inactivation and endothelial cell preservation of Merk expression in the brain (29). When we injected corn oil control into APP/PS1Cx3cr1\(^{CreER}\)Merk~\(^{f/f}\) mice immediately after weaning (at P30 to 40), we observed an onset and progression of sudden death that closely paralleled that seen in APP/PS1 (SI Appendix, Fig. S1). In contrast, when we injected tamoxifen into these same mice, we observed a sudden death time course that paralleled that seen in APP/PS1Merk~\(^{-}\)/C0 mice (SI Appendix, Fig. S1). Although brain (and other) perivascular macrophages are also Cx3cr1~\(^{-}\) and weakly express Mer (35), these cells account for <3% of the CD45~\(^{+}\) cells in the adult brain and are vastly outnumbered by Mer~\(^{+}\) microglia (36). We therefore conclude that the cells that drive the sudden death in the germline APP/PS1Merk~\(^{-}\) mutants are microglia. These brain macrophages respond to and regulate neuronal activity (37, 38) and are strongly activated, especially in the hippocampus, subsequent to seizures (39–41). As discussed below, they are also exceptionally active TAM-dependent phagocytes (29).

The major contribution of Mer mutation and the minor contribution of Axl mutation to enhanced early lethality in the
APP/PS1 compound mutants (Fig. 1) are both consistent with the fact that microglia normally express high levels of Mer and low levels of Axl (22, 29). (In contrast, perivascular macrophages express high levels of Axl and low levels of Mer) (35). Although microglial Axl is markedly up-regulated in APP/PS1 and other mouse AD lines (22, 36) and in human AD (42), this up-regulation occurs exclusively in microglia that are tightly bound to Aβ plaques (22, 43). The first deposition of Aβ plaques in the cortex and hippocampus of APP/PS1 mouse is observed at ~P120, and plaques are not reliably detected across the brain in these mice until ~P150 (44, 45). Thus, nearly all of the sudden death quantified in Fig. 1 occurred before the initial deposition of Aβ plaques and the up-regulation of Axl. Indeed, as detailed below, sudden death and the development of plaques were strongly anticorrelated in all genotypes. Although the contribution of Axl mutation to enhanced APP/PS1 lethality was minimal, most comparative analyses below were performed with Axl−/−Mertk−/− compound mutants to ensure complete TAM inactivation.

**APP/PS1Mertk−/− Seizures Are Associated with Massive Firing of the Dentate Gyrus.** As noted above, prior analyses demonstrated that young APP/PS1 mice display repeated episodes of cortical epileptiform seizure activity and experience tonic–clonic seizures that very frequently lead to acute sudden death (12, 13). These events result from the overproduction of Aβ (19), which triggers cortical and hippocampal hyperexcitability via sustained elevation of the resting membrane potential of layer 2/3 pyramidal neurons and dentate granule cells, respectively (12). As also noted above, seizures are a recognized comorbidity at early stages of human AD (46, 47). Continuous video monitoring of APP/PS1Axl−/−Mertk−/− and APP/PS1Mertk−/− mice indicated that sudden death in these compound mutants was, consistent with the previous reports discussed above, also the result of violent tonic–clonic seizures (SI Appendix, Movies S1 and S2 and Fig. S2A).

The V-shaped DG of the hippocampus normally displays low spiking activity and is thought to function as an inhibitory gate that limits excitatory input to the remainder of the hippocampus. It is a common initiating locus for temporal lobe epilepsy in humans and mice (48–50), when this gating is weakened and granule cells become hyperexcitable. We did not perform any manipulation to induce seizures in our colonies and perform any manipulation to induce seizures in our colonies and are normally encountered four APP/PS1Mertk−/− mice during or immediately after a spontaneous lethal seizure. Although brains from these long-dead animals could not be analyzed (decomposition begins within minutes of death), we fortuitously encountered four APP/PS1Mertk−/− mice that had died overnight. Although brains from these long-dead animals could not be analyzed (decomposition begins within minutes of death), we fortuitously encountered four APP/PS1Mertk−/− mice during or immediately after a spontaneous lethal seizure. Immunostaining for the transcription factor Arc (Arg3.1), an established marker of neuronal activity (51), demonstrated massive firing of all of the granule cells in the DG in these mice (Fig. 2A, Top). (In contrast to our normal immunohistochemical analyses, which were performed on sections cut from perfusion-fixed brains, this Arc immunostaining was performed on sections cut from immersion-fixed brains removed shortly after death. All representative panels from both of the genotypes in Fig. 2 and SI Appendix, Fig. S2 are from sections of immersion-fixed material.) In addition to the DG, widespread high Arc staining was evident in NeuN+ neurons across many loci of the APP/PS1Mertk−/− brain after seizure, including the CA3 field of the hippocampus, cortex, striatum, hypothalamus, superior colliculus, and olfactory bulb (Fig. 2A, Top and SI Appendix, Fig. S2B).

This widespread Arc expression stood in marked contrast to that seen in mice of the same genotype in the absence of seizure (Fig. 2A, Middle). In these animals, Arc+ loci of neuronal firing were still evident in the cortex (SI Appendix, Fig. S2C), suggestive of a basal level of network hyperactivation, but the massive activation of all neurons in the DG was not seen. Instead, only occasional DG cells were Arc+ (Fig. 2A, Middle Right), consistent with the normally sparse Arc expression and firing of DG granule cells that has been documented for wild-type mice (50, 52, 53). This same sparse expression of Arc in the DG was also seen in Mertk−/− mice (Fig. 2A, Bottom).
Importantly with respect to the results presented below, these Merik<sup>f/f</sup> mice displayed only minimal Arc expression in the cortex (Fig. 2 A, Bottom; compare with seizure-free APP/PS1Merik<sup>−/−</sup> mice in SI Appendix, Fig. S2C), and never developed seizures. Quantification revealed that >90% of NeuN<sup>+</sup> neurons were Arc<sup>+</sup> in the cortex, DG, CA3, and striatum of APP/PS1Merik<sup>−/−</sup> mice after lethal seizure (Fig. 2B). This fraction was ≤10% in the striatum and DG of the same APP/PS1Merik<sup>−/−</sup> mice in the absence of seizure, but consistent with the immunostaining in SI Appendix, Fig. S2C, was ∼35% in the cortex (Fig. 2B). The fraction of NeuN<sup>+</sup> neurons that were Arc<sup>+</sup> was ≤10% in all brain regions for Merik<sup>−/−</sup> mice in the absence of seizure (Fig. 2B).

**TAM Mutation Leads to the Accumulation of Apoptotic Cells in the DG.** Together with the SVZ adjacent to the lateral ventricle, the DG of the hippocampus is a significant site of young adult neurogenesis in the mouse brain (27, 54). DG neurogenesis occurs during the first months of postnatal life, in a stem cell population that resides in its subgranular zone (SGZ). This SGZ is a thin layer of cells situated immediately beneath (interior to) the mature granular cell layer (GCL) (55), which is marked by very high cell density. (Dotted lines in Fig. 3A panels delimit the center of the GCL in the DG.) In the SGZ, it has been observed that ∼80% of newborn cells never survive to differentiate into DG granule cell neurons, but instead die by apoptosis within 10 d of their birth (56). These ACs are normally phagocytically cleared exclusively by microglia (56), and we have previously shown that microglia, both in the SGZ and the SVZ, require TAM receptors to effectively carry out AC clearance (28, 29). In the SGZ, for example, we have shown that Axl<sup>−/−</sup>Merik<sup>−/−</sup> mice display marked reductions in phagocytic index and capacity and a 2-fold increase in AC density and that tamoxifen-injected Cx3cr1<sup>CreER<sub>+</sub>Merik<sup>−/−</sup></sup> microglial-specific Mer knockouts (29) also display reductions in phagocytic index and capacity and a 1.6-fold increase in AC density (28). Similarly, we have previously shown that the dramatic accumulation of ACs seen in the Axl<sup>−/−</sup>Merik<sup>−/−</sup>SVZ is substantially phenocopied in the SVZ of tamoxifen-injected Cx3cr1<sup>CreER<sub>+</sub>Merik<sup>−/−</sup></sup> mice (29).

We used immunostaining for cleaved caspase 3 (cCasp3) to identify ACs, together with the lysosomal marker CD68 to monitor microglial activation, in the hippocampus of WT, APP/PS1, Axl<sup>−/−</sup>Merik<sup>−/−</sup>, and APP/PS1Axl<sup>−/−</sub>Mertk<sup>−/−</sup> mice (Fig. 3A). We performed this analysis at P30, near the onset of sudden death in all genotypes (Fig. 1). We found that while the density of cCasp3<sup>+</sup> ACs in the DG exhibited a nonsignificant increase in APP/PS1 mice relative to WT, the combination of APP/PS1 with Axl<sup>−/−</sub>Mertk<sup>−/−</sup> double mutation resulted in a remarkable 6-fold increase in the steady-state level of these dead cells (Fig. 3 A and B). This is in contrast to the 2-fold increase in AC density seen in Axl<sup>−/−</sup>Merik<sup>−/−</sup> alone. The area occupied by CD68<sup>+</sup> microglial puncta was increased 3.3-fold in the APP/PS1Axl<sup>−/−</sub>Mertk<sup>−/−</sup> DG relative to WT (Fig. 3 A and B). Essentially all CD68<sup>+</sup> cells were Iba1<sup>+</sup> microglia (SI Appendix, Fig. S3A), and consistent with our earlier work (29), the Axl<sup>−/−</sub>Mertk<sup>−/−</sup> double mutation led to a marked increase in microglial Iba1 expression (SI Appendix, Fig. S3B). As has been described previously (29), AC accumulation was exclusively seen in Axl<sup>−/−</sub>Mertk<sup>−/−</sup>-neurogenic regions and was undetectable elsewhere in the Axl<sup>−/−</sub>Mertk<sup>−/−</sup>-brain. Consistent with an extensive prior literature (57) and our own analyses in the SVZ (29), these ACs were NeuN<sup>−</sup> but expressed the neural stem cell marker doublecortin (DCX) at both P30 and P90 (SI Appendix, Fig. S3D).

**Fig. 3.** Marked accumulation of apoptotic cells and activation of microglia in the P30 APP/PS1Merik<sup>−/−</sup>-dentate gyrus. (A) Representative sections through the DG of mice of the indicated genotypes stained for nuclei (Hoechst, blue, Top), the microglial activation marker CD68 (white, Middle), and the apoptotic cell marker cleaved Caspase-3 (cCasp3, green, Lower). The SGZ, where adult neurogenesis occurs, is arrowed, and the center of the GCL of the DG is marked by dotted lines. Note that most apoptotic cells in APP/PS1 Axl<sup>−/−</sub>Mertk<sup>−/−</sup> mice accumulate in the SGZ (Lower Right). (B) Quantification of the results represented in A. Data in B and all subsequent figures in the paper are represented as mean ± 1 STD. Data are collected from five serial hippocampal sections per animal and n = 3 to 6 per genotype. One-way ANOVA test. (Scale bar, 100 μm.)
We therefore performed similar BrdU pulse labelings in APP/PS1 versus APP/PS1Axl−/−Mertk−/− mice at either P30, when hippocampal neurogenesis is robust, or at P90, when neurogenesis is reduced (see below). We then quantified labeled cells at 28 and 7 d after the pulse. As illustrated in the representative example shown in Fig. 4A, we counted BrdU+ cells in both the SGZ and GCL and also in the entire DG (including the hilus and molecular layer). In the P90 series, we also stained sections for the proliferation marker Ki67 (Fig. 4A). As noted above, hippocampal neurogenic cell division occurs specifically in the SGZ, from which surviving cells subsequently migrate into the GCL and differentiate into neurons. At 28 d after a BrdU pulse applied at P30, we measured a ~2.1-fold increase in labeled cells in APP/PS1Axl−/−Mertk−/− versus APP/PS1 mice in both the SGZ + GCL and the full DG quantifications (Fig. 4 B and C). Note that, as expected, the large majority of BrdU+ cells were located in the SGZ and GCL (Fig. 4 B and C). By 28 d after a BrdU pulse, most of the labeled cells that subsequently undergo apoptosis, together with live labeled cells that are engulfed by phagoptosis, have been cleared by microglial phagocytosis (28). These processes are not complete by 7 d after a pulse, and so we detected more labeled cells overall at 7 d after a BrdU pulse into P90 mice (Fig. 4 D and E). (Again at P90, essentially all BrdU+ cells were present in the GCL and SGZ; SI Appendix, Fig. S3E). Correspondingly, we quantified an even greater disparity between APP/PS1Axl−/−Mertk−/− and APP/PS1 mice in these P90 experiments and measured ~3.5-fold more BrdU+ or BrdU+Ki67+ cells in the former relative to the latter. Together, these results indicate that introducing TAM mutations into the APP/PS1 line leads to the exuberant introduction of new cells into the GCL of the DG during the period of adult neurogenesis.

TAM Mutation Leads to Extra Mossy Fiber Synapses in CA3. All of the cells that survive and differentiate as a result of adult neurogenesis in the SGZ are excitatory glutamatergic neurons that project mossy fibers from the GCL to neurons in the hilus and the CA3 field of the hippocampus (63). We therefore asked whether the enhanced survival of newborn cells in the APP/PS1Axl−/−Mertk−/− GCL resulted in more excitatory connections between the DG and CA3, leading to an excitatory/inhibitory imbalance. To address this question, we immunostained sections of the P30 APP/PS1 and APP/PS1Axl−/−Mertk−/− hippocampus with an antibody to the ZnT3 vesicular zinc transporter, which is expressed in the synaptic vesicles of GCL neurons and has previously been shown to intensely and specifically label presynaptic mossy fiber terminals in the hilus and CA3 (64, 65) (Fig. 5A). Hippocampal CA3 also receives excitatory glutamatergic input from entorhinal cortex and CA3 collaterals, but by using ZnT3 and focusing on CA3 proximal dendrites, we have almost exclusively assessed mossy fiber synapses. We simultaneously immunostained the same hippocampal sections with an antibody to PSD-95, which labels postsynaptic densities. We then used Airyscan superresolution imaging of the proximal region of CA3 pyramidal dendrites (box in Fig. 5A, enlarged in Fig. 5B), and Imaris software to construct ZnT3 and PSD-95 spheres (Fig. 5 C) for synaptic quantification by proximity measurement of red (ZnT3) and

![Fig. 4.](image-url) Excess newborn cells in the APP/PS1 and APP/PS1Axl−/−Mertk−/− dentate gyrus. (A) Representative panels of the same sagittal sections through the V-shaped DG of APP/PS1 (left panels) and APP/PS1Axl−/−Mertk−/− mice (right panels) in which dividing cells were pulse labeled with BrdU at P90 and then analyzed 7 d later. Top row, cell nuclei labeled with the DNA dye Hoechst 33258; second row, BrdU-labeled nuclei visualized with an anti-BrdU antibody; third row, cells positive for the proliferation marker Ki67 visualized with an anti-Ki67 antibody; and bottom row, merged images. Relevant hippocampal regions indicated in the top panels are ML, molecular layer; GCL, granule cell layer; SGZ, subgranular zone; CA3, cornu ammonis field 3. Dotted lines indicate the center of the GCL (white) and the thin layer of the SGZ (red). (B and C) Quantification of BrdU+ cells in the indicated regions of APP/PS1 and APP/PS1Axl−/−Mertk−/− mice pulse labeled at P90 and analyzed 28 d later. (D and E) Quantification of BrdU+ cells in the indicated regions of APP/PS1 and APP/PS1Axl−/−Mertk−/− mice pulse labeled at P90 and analyzed 7 d later. Data are average from five serial hippocampal sections per animal and n = 3 to 4 per genotype. Unpaired t test. (Scale bars, 200 μm.)
green (PSD-95) spheres (Method). As is standard in the field, we defined synapses as ZnT3/PSD-95 sphere pairs that were ≤0.7 μm apart.

These analyses revealed that at 1 mo after birth, the CA3 proximal dendrite area of the APP/PS1Axl/Merk−/− hippocampus exhibits ~38% more mossy fiber synapses than are seen in APP/PS1 at the same age, as defined by ZnT3/PSD-95 coincidence (Fig. 5D). They are consistent with the neuronal differentiation of a subset of the excess BrdU+ cells seen in the APP/PS1Axl/Merk−/− hippocampus 28 d after a pulse labeling (Fig. 4), and with the subsequent development of neuronal connections between these GCL neurons and their synaptic targets in CA3.

Sudden Death Is Coincident with Neurogenesis but Anticorrelated with Plaques. The level of DG neurogenesis, which is largely restricted to young adulthood in both mice and humans, has previously been precisely measured over the first 9 mo of life in C57BL/6J mice (66). This is the strain onto which all of the mutations in Fig. 1 have been backcrossed and maintained. We therefore compared the time course of sudden death in APP/PS1Axl/Merk−/− mice and APP/PS1Merk−/− mice after lethal seizure in APP/PS1Axl/Merk−/− mice (Fig. 1) to the time course of expression of Ki67 (Fig. 5), and with the subsequent development of neuronal connections between these GCL neurons and their synaptic targets in CA3.

At the same time that sudden death and hippocampal neurogenesis were coincident, we observed that sudden death and the development of dense-core amyloid plaques in all genotypes were strongly anticorrelated. We (and others) have previously quantified the development of dense-core Aβ plaques, as identified by staining with the amyloid dye ThioS, in APP/PS1 mice over time (22). Our earlier analyses of cortical and hippocampal dense-core plaque burden began at 7 mo (22), after essentially all sudden death in APP/PS1Axl/Merk−/−, APP/PS1Merk−/−, and APP/PS1Gas6−/− mice had already occurred (Fig. 1). We therefore extended our quantification of ThioS Aβ plaque burden to earlier time points and compared plaque development to sudden death over time. We also included previously published sudden death data from APP/PS1 mice lacking the interleukin-18 receptor, which similarly display enhanced early lethality relative to APP/PS1 alone (ref 13). These analyses demonstrated that the cessation of early death in the APP/PS1Axl/Merk−/− mice coincided with the first detectable deposition of ThioS dense-core Aβ plaques (SI Appendix, Fig. S4C). No further sudden death was observed over the subsequent course of plaque accumulation in any line.

Discussion

Many investigators working with APP/PS1 mice and other amyloidogenic mouse AD models have heretofore noted the occurrence of early sudden death in their colonies (12, 13, 17, 19, 22). The strong association between seizures and early AD notwithstanding, this early lethality has for the most part simply been ignored, since it occurs before the development of amyloid plaques and results in the loss of a relatively small number of experimental animals. At the same time, seizures early in the course of human AD are almost never fatal. The increase in early lethality upon the introduction of TAM system mutations into APP/PS1 mice documented above is something that we of course observed during our previously published analyses of these mice (22), since it severely limited the number of animals that were available for study. We did not discuss this phenomenon at the time because our work was limited to a dissection of the role of TAM signaling in the microglia that surround Aβ plaques, and as shown above, the exacerbation of early APP/PS1 lethality by TAM mutation occurs, just as in APP/PS1 alone, before any plaques appear. In addition, we didn’t have a mechanistic explanation for the phenomenon.

The results documented above indicate that this explanation involves an increased excitatory drive from the DG to CA3 target neurons in the hippocampus. We found that the DG was the single most heavily activated region of the brain in TAM-deficient APP/PS1 mice after seizure. We further found that TAM system mutations allow for the survival and subsequent differentiation of an excess number of excitatory granule cell neurons in the DG—viable cells that would normally be cleared by TAM-dependent microglial phagocytosis (phagoptosis). This is essentially the same phenomenon and the same magnitude of effect that we have previously documented in the other center of adult neurogenesis in the mouse brain—the SVZ—where mutation of Axl and Merk disables microglial phagocytosis and leads to the excess accumulation of viable newborn neurons in the olfactory bulb (29). The added cells that we detect in the DG may form additional synapses onto CA3 neurons, providing a hippocampal excitatory/inhibitory imbalance that feeds into the existing cortical hyperexcitability of APP/PS1 mice. This ignites widespread firing across many regions of the TAM-deficient APP/PS1 brain. It is noteworthy that after lethal seizure in APP/PS1Merk−/− mice, the only hippocampal field other than the DG to display significant activation, as marked by elevated Arc expression, is CA3.

This hippocampal field is situated at a crucial location and plays an important role in the encoding of spatial representations and memory. Mossy fiber inputs from the DG pass highly integrated and sparsely coded cortical inputs on to CA3. As
such, this pathway and its synaptic connections are exceptionally susceptible to both seizures and the neurodegeneration that is exacerbated by Aβ-induced neuronal hyperactivity (67–69). AD patients with epileptic activity and seizures have been reported to have impaired cortical–hippocampal circuits and hastened cognitive decline (70–72); and in line with preclinical studies (18, 73), a recent phase IIa human clinical trial demonstrated that low-dose treatments with the anticonvulsant drug levetiracetam can improve deficits in hippocampal-mediated spatial memory and executive function specifically in AD patients who experience seizures and present with network hyperactivity (74).

Since we argue that the greatly enhanced lethality seen in APP/PS1AxlflMertkfl mice relative to APP/PS1 alone is due to compromised microglial phagocytosis, similar phenotypes might be predicted to develop in an amyloidogenic mouse AD model if phagocytic microglia were simply depleted from such a model before birth. An analysis of this form has very recently been reported (75). Kiani Shabastari et al. exploited a mouse line in which an essential enhancer element in the mouse colony stimulating factor 1 receptor (Csf1r) gene was deleted (76). Because microglia and many other tissue macrophages require continuous signaling through the Csf1r receptor for their survival (77, 78), deletion of this Csf1r enhancer element results in the embryonic ablation of microglia (76). When these Csf1r mutants lacking microglia (designated “FIRE mice”) were crossed into the 5xFAD amyloidogenic AD model, it was observed that >50% of the compound mutants died from sudden death, and that most of this death occurred before P150 (75). The correspondence between this phenotype and the phenotype we describe above is striking. Like APP/PS1 mice, 5xFAD mice exhibit a low incidence of sudden death before P150, and like AxlflMertkfl mice (or Mertkfl or Gas6fl mice), FIRE mice display no lethality on their own (75). It is only when combined with the amyloidogenic APP/PS1 and 5xFAD alleles that the sudden death contributions of the TAM system and FIRE mutations are revealed. Although Kiani Shabastari et al. did not provide a mechanistic explanation for the profound increase in early death seen when the FIRE enhancer mutation is introduced into the 5xFAD model (75), we propose that their death phenotype and ours are similar because both are tied to a deficit in microglial phagocytosis in the hippocampus.

Greatly enhanced early mortality, again due entirely to lethal seizures, has also been seen when the APP/PS1 line is crossed with knockouts for the inflammatory cytokine interleukin-18 (IL-18) or its receptor (13). Cortical hyperexcitability and early lethal seizures in APP/PS1IL18fl mice, which also can be effectively suppressed by levetiracetam, are associated with enhanced expression of excitatory synaptic proteins in the hippocampus, marked perturbations in hippocampal physiology, and increased sensitivity to the seizures that are chemically induced with picrotoxin (13). Anomalies in adult neurogenesis in the SGZ and DG similar to those that we have detected in AxlflMertkfl mice, including an increased number of BrdU+ cells in the DG (relative to WT) at 24 h after a BrdU pulse, have also been reported for IL18fl mice (79). With respect to these IL-18 effects, it is of interest to note that, within the brain, the only cells that express the full complement of NLRP3 components required to transduce IL-18 inflammatory signals are microglia. Since IL-18 receptors are predominantly expressed by neurons (80), a role for microglial IL-18 in the suppression of excitatory neurotransmission has also been advanced (13). Our results suggest that TAM-dependent microglial phagocytosis similarly suppresses the elaboration of exuberant excitatory connections between DG neurons and their targets in CA3.

Although mutations in the Mertk and Gas6 genes dramatically increase the incidence of lethal seizures when crossed into APP/PS1, it is important to emphasize that these mutations only function as “second hits” (this is also the case for IL-18 and IL-18R mutations when introduced into APP/PS1 and for the FIRE mutation when introduced into 5xFAD) (13, 75). Seizure-induced sudden death is routinely observed in the APP/PS1 line alone, but never in Mertkfl, Axlfl, Gas6fl, or AxlflMertkfl mice. Thus, the key triggering event with respect to seizure induction in all cases is the production of excess Aβ from the mutant human transgenes present in the APP/PS1 locus: The additional excitatory glutamatergic connections that we detect between the DG and CA3 are not on their own sufficient to trigger seizures. As previously described, excess Aβ is associated with sustained cortical and hippocampal hyperexcitability (12).

Why lethal seizures are not observed after ~P150 in APP/PS1 lines, with or without TAM mutations, remains a mystery. However, the observation that these seizures stop when dense-core plaques first appear is intriguing. It has previously been shown that the application of Aβ protofibrils to neurons in slice cultures of mouse cortex and hippocampus results in membrane depolarization (resting membrane potential elevation) and increased neuronal excitability (12). Recent work has shown that these protofibrils are normally phagocytosed by microglia, where they are routed to microglial lysosomes and then concentrated and aggregated (21, 77, 81, 82). This highly aggregated Aβ is finally deposited into the dense cores of dense-core plaques (81), and without microglia and microglial phagocytosis it now appears that these structures cannot be formed (77). As such, it seems possible that the cessation of seizures in APP/PS1 mice may be due in part to the removal of seizure-inducing forms of Aβ polymers by microglial phagocytosis and their sequestration into dense-core plaques. The TAM RTKs are currently being therapeutically targeted—using antibody inhibitors, ligand traps, and small-molecule inhibitors of the TAM kinases—in the context of cancer, since the growth and metastasis of many different cancers are promoted by TAM up-regulation (83–85). The adverse effects that these potential therapeutics might have on microglial phagocytosis in the brain have not been considered.

Methods

Mice. The Axlfl (23), Mertkfl (23), AxlflMertkfl (23), Gas6fl (32), Cx3cr1fl (34), and Mertkfl (29) mouse strains have been described previously. (Contact Carla Rothlin@yale.edu for Mertkfl). B6.Cg-Tg(APPswPSEN1c99Tg) hemizygous mice (20, 21) (APP/PS1) (JAX No. 005864) were crossed with Axlfl, Mertkfl, AxlflMertkfl, Gas6fl, and/or Cx3cr1fl lines. Recombination (inactivation) of the Mertkfl allele in Cx3cr1flCotahMertkfl mice was achieved using tamoxifen injection at weaning at P28 to 30. APP/PS1 Cx3cr1flCotahMertkfl mice received two doses (150 mg/kg body weight) of tamoxifen dissolved in 10% EtOH 90% filtered corn oil (Sigma) by intraperitoneal (IP) injection with the two successive injections 48 h apart. Control mice received two successive IP injections of vehicle (corn oil) alone. Mice were analyzed for Mer expression 1 wk, 2 mo, and 7 mo after the injection to ensure efficient and long-lasting Mer inactivation. All lines have been backcrossed for >10 generations to and maintained on a C57BL/6 background. All animal procedures were conducted according to protocols approved by the Salk Institute Animal Care and Use Committee (IACUC) (protocol No. 17.00009). Mice of both genders were randomly allocated to experimental groups unless otherwise noted.
Reagents and Antibodies. BrdU was from Sigma-Aldrich. Antibodies used were as follows: anti-mouse Ad (R&D AF8544), anti-Mer (eBioscience D55MM and R&D AF5911), anti-mouse Gas6 (R&D AF8966), anti-BrdU (BU1/75 [ICR1], AbD serotec), anti-ib1 (Wako 019-19741, 1,200, and Novus NB100-1028), anti-cleaved Casp3 (Asp175, Cell Signaling 9661), anti-doublecortin (DCX, Millipore ab2253), anti-ZnT3 (Syss 197 002, anti-PDSD-95 (Abcam ab18258), anti-CD68 (BioRad MCA1957), anti-NeuN (Chemicon MAB377), anti-Arc (Syss #156003), anti-kc6 (Abcam ab15580), and anti-kc6-APC-conjugated (Invitrogen, clone ScAl51). Secondary antibodies for immunohistochemistry were fluorophore-conjugated anti-rat (712-545-153 and 712-155-153 from Jackson ImmunoResearch), anti-guinea pig (Molecular Probe A-11073), and anti-rabbit (A-10040 or A-21206 from Life Technologies and 711-606-152 from Jackson ImmunoResearch), and anti-mouse (A-21202 from Molecular Probe, 715-165-150 and 715-175-150 from Jackson ImmunoResearch). Key reagents and their usage are as follows: ThioS (Acros Organics 213150250), Hoechst 33258 in a 1% solution of bisbenzimide in water (Sigma-Aldrich B-2883), and paraformaldehyde (PFA) (Sigma P6148).

BrdU Pulse Labeling. Mice were administered a single dose of BrdU (100 mg/kg, dissolved in Dulbecco’s phosphate-buffered saline (D-PBS)) intraperitoneally at P30 or P90 (28) and returned to their home cage. At 7 d or 28 d after BrdU injection, mice were anesthetized and their brains extracted using routine perfusion procedures as described previously (29).

Immunohistochemistry. Mice were anesthetized with a final concentration of 100 mg/kg ketamine/10 mg/kg xylazine in accordance with IACUC guidelines and were then transcardially perfused with 20 IU/ml heparin in PBS followed by freshly prepared 4% PFA in PBS. Both brain hemispheres were postfixed in 4% PFA in PBS overnight and subsequently infiltrated in 30% sucrose in PBS for 1 d and flash frozen in Tris-buffer saline (TBS) tissue freezing medium (TFM) (General Data Healthcare TFM-5) using isopentane and kept at 80°C until use. Frozen brain hemispheres were sagittally cryosectioned at 15 μm, air-dried overnight, and subsequently processed for staining. Antigen retrieval with citrate buffer was accomplished either by heating sections at 100°C for 3 min in prewarmed citrate buffer (pH 6.0) or by 1 h incubation in blocking buffer (PBS containing 0.3% Triton X-100, 0.1% Tween-20, 2% donkey serum and 2% IgG-free bovine serum albumin (BSA)). Sections were then incubated overnight at 4°C with primary antibody diluted in blocking buffer, then washed in PBS 0.1% Tween-20 and incubated for 2 h at 22°C in the dark with fluorophore-coupled secondary antibodies and subsequently in Hoechst for 5 min diluted in blocking buffer. Sections were washed, sealed with Fluoromount-G (SouthernBiotech), and stored at 4°C. For immunofluorescence experiments, brains from mice found dead immediately after seizure or control mice after receiving anesthesia until respiration ceased were freshly extracted after decapitation, split down the midline, and immediately immersion fixed in freshly prepared 4% PFA in PBS for 48 h. Brains were subsequently processed as for routinely perfused brains.

Dense-Core Plaque Labeling and Quantification. Cortical and hippocampal plaque density across the course of disease at the indicated timepoints was quantified by staining with the fluorescent amyloid dye thioflavin S (ThioS) as described previously (22). ThioS binds to the β-pleated sheets of dense-core amyloid plaques and provides for reliable semiautomated quantification of plaques. ThioS plaque density quantification was performed as described previously (22). Briefly, a serial set of 15 μm brain sagittal sections (six sections/mouse) was obtained 0.25 mm to 0.85 mm from the midline, each spaced 0.15 mm apart, and mounted directly on glass slides after cryosectioning. Serially cut sections from both hemispheres and various timepoints were used for measuring plaque density by histochemistry with ThioS staining using published protocols (86). Briefly, slides with brain sections were air-dried, then sequentially dehydrated in 70% and 80% ethanol before staining in filtered 1% ThioS dissolved in 80% ethanol in the dark. Slides were then rehydrated and washed sequentially in 80% ethanol, 70% ethanol, and water before being counterstained with Hoechst. ThioS-labeled brain sections were then either directly mounted or followed by further immunofluorescence staining, stored at 4°C, and imaged within a week. Image analysis of the number of cortical and hippocampal plaque density quantification was done blind with respect to genotypes and in a semiautomated fashion with set intensity thresholds for all conditions, using the “analyze particles” function in ImageJ or Fiji software.

Survival and Home-Cage Video Monitoring. Long-term (2 wk) continuous digital videos of young adult AD mice (1.5 to 3 mo) housed in their home cages with clear cage lids (two mice per cage) were collected at 25 fps using a high-resolution, infrared-equipped, digital video camera (Sony HandyCam HDR-XR520), set up in a top-down fixed angle for recording. The recording room was maintained on a 12-h/12-h light/dark light cycle. Video monitoring of sporadic deaths and spontaneous convulsive seizures was performed by reviewing video footage via a VLC player. Cages were checked every 12 h to ensure proper air conduction, ample and accessible food and water, and to collect any mouse found dead. Brains from mice found dead during the study were freshly extracted, immersion fixed, and immunostained for Arc expression.

Confocal Microscopy. Fluorescent immunostaining images were acquired with a Zeiss LSM 710 confocal microscope using Plan-Apochromat 20× 0.8 NA air-matched or 63× 1.4 NA oil objectives (laser lines: 405 nm, 488 nm, 594 nm, and 633 nm). Image size was 1,024 × 1,024 pixels. For quantification of Arc+/NeuN+ cells in various brain regions, two representative single-plane images, at 320 μm × 320 μm per section from two sections per biological sample, were randomly taken from striatum, DG, CA3, or layers 3 to 4 of prefrontal cortex using Hoechst as an unbiased channel. For synaptic analysis, Aiyrsan superresolution images were acquired with a Zeiss LSM 880 Rear Port Laser Scanning Confocal and Aiyrsan FAST Microscope using 63× oil-immersion objective and piezo z-stage for superresolution imaging. Three images from five serial brain sections per animal containing CA3 apical dendrites were imaged as 8-bit images at 2,124 pixels × 2,124 pixels and 53.98 μm × 53.98 μm as a z stack of seven slices with a total thickness of 1.29 μm. Exposure acquisition was set according to the nonmutant samples and all sections were imaged within the same session. Images were obtained and processed via the Zen Black edition. Images for cortical and hippocampal plaque quantification were acquired with an Olympus VS-120 Virtual Slide Scanning Microscope.

Data Analysis. For fixed-brain thin sections, maximum intensity projection images of 212 μm × 212 μm (1,024 × 1,024 pixel resolution) were analyzed in Fiji. Approximately five to seven plaques per brain section (three to five sections per animal) were randomly chosen in the prefrontal cortex using the 6610 channel. For analyses in the P30 DG, tiled and stitched maximum intensity projection images from field of view of 640 μm × 1,280 μm (1,024 × 1,024 pixel resolution) that fully captured the DG were used for cCasp3 number and area analysis using particle analysis in Fiji with a 3 μm2 minimum cutoff. Higher magnification images 212 μm × 212 μm (1,024 × 1,024 pixel resolution), which centers on dentate gyrus/SZG were used for CD68 area analysis in P30 DG. For adult neurogenesis analysis in hippocampus using BrdU, the analysis of neurogenesis in vivo was performed as described previously (28, 87). Five sagittal sections from one series per animal of each genotype containing the same area of hippocampus were analyzed in all experiments. Proliferating cells were assessed by BrdU+ and K-/67+ cell quantification and neuroblasts were assessed by Dcx+ cell quantification, while mature neurons were assessed by NeuN+ cell quantification. The proliferation of either of these populations was quantified by their colocalization with BrdU or K-/67 using the software ImageJ (Fiji). Analysis of synapse number was performed blind to genotype using IMARIS software (version 9.1.2, bitplane). Synapses were assessed as proximity of the postsynaptic PSD-95 puncta to presynaptic ZnT3 puncta. The z stacks of each condition (APP/PS1 or APP/PS1 Ad+/Merk−/−) were presented as three-dimensional volumes acquired by Aiyrsan superresolution imaging. Puncta from each channel were defined in each channel as spheres of average diameter of 0.5 μm using the spot detection function. The localization of spots from different channels within a distance from center to center of spots of 0.7 μm was quantified as a synapse. Synapse number from each image was averaged across five sections per animal.

Statistical Analysis. Statistics were performed with GraphPad Prism (version 8.0) software.

Data, Materials, and Software Availability. All study data are included in the article and/or supporting information.
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