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ORIGINAL PAPER



Deficiency of TYROBP, an adapter protein for TREM2 and CR3 receptors, is neuroprotective in a mouse model of early Alzheimer's pathology

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Abstract Conventional genetic approaches and computational strategies have converged on immune-inflammatory pathways as key events in the pathogenesis of late onset sporadic Alzheimer's disease (LOAD). Mutations and/ or differential expression of microglial specific receptors such as TREM2, CD33, and CR3 have been associated with strong increased risk for developing Alzheimer's disease (AD). DAP12 (DNAX-activating protein 12)/ TYROBP, a molecule localized to microglia, is a direct partner/adapter for TREM2, CD33, and CR3. We and

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others have previously shown that TYROBP expression is increased in AD patients and in mouse models. Moreover, missense mutations in the coding region of TYROBP have recently been identified in some AD patients. These lines of evidence, along with computational analysis of LOAD brain gene expression, point to DAP12/TYROBP as a potential hub or driver protein in the pathogenesis of AD. Using a comprehensive panel of biochemical, physiological, behavioral, and transcriptomic assays, we evaluated in a mouse model the role of TYROBP in early stage AD. We crossed an Alzheimer's model mutant APPKM670/671NL/ $PSEN1^{\Delta exon9}$ (APP/PSEN1) mouse model with Tyrobp^{-/-} mice to generate AD model mice deficient or null for $Tyrobp^{+/-}$ or *APP/PSEN1*; TYROBP (APP/PSEN1; *Tyrobp*^{-/-}). While we observed relatively minor effects of TYROBP deficiency on steady-state levels of amyloid-β peptides, there was an effect of Tyrobp deficiency on the morphology of amyloid deposits resembling that reported by others for $Trem2^{-/-}$ mice. We identified modulatory effects of TYROBP deficiency on the level of phosphorylation of TAU that was accompanied by a reduction in the severity of neuritic dystrophy. TYROBP deficiency also altered the expression of several AD related genes, including Cd33. Electrophysiological abnormalities and learning behavior deficits associated with APP/PSEN1 transgenes were greatly attenuated on a Tyrobp-null background. Some modulatory effects of TYROBP on Alzheimer'srelated genes were only apparent on a background of mice with cerebral amyloidosis due to overexpression of mutant APP/PSEN1. These results suggest that reduction of TYROBP gene expression and/or protein levels could represent an immune-inflammatory therapeutic opportunity for modulating early stage LOAD, potentially leading to slowing or arresting the progression to full-blown clinical and pathological LOAD.

Keywords TYROBP/DAP12 · TREM2 adapter · CR3 adapter · Alzheimer's disease · Microglia · APP/PSEN1

Introduction

Conventional wisdom has held that the chronic neuroinflammation associated with LOAD may be a secondary or even protective event that occurs in response to AB deposition and may occur only in late stages of AD. However, recent genetic and genomic approaches, as well as computational strategies, have converged on immune-inflammatory pathways as risk factors and as key events in the pathogenesis of late-onset sporadic Alzheimer's disease (LOAD) [19]. Moreover, correlation between inflammatory genes and clinical presentation of previously asymptomatic cerebral amyloidosis (ACA) indicates a role for inflammation and microglia in the progression from ACA to the earliest stages of mild cognitive impairment (MCI) and/or mild clinical AD. Among the genes implicated by the largest available genome-wide association studies [43], one-third is either unique to, or enriched in, microglia. Recently identified mutations and variants in genes encoding important immune receptors including CD33, CR3 (Complement Receptor 3), and TREM2 (Triggering Receptor Expressed On Myeloid Cells 2), have been genetically linked to LOAD risk, highlighting the potential role of a dysregulated immune response in an early, and perhaps causative role in AD pathogenesis. Unlike autosomal dominant familial Alzheimer's mutations that promote elevation of the $A\beta 42:40$ ratio or of other variant hyperaggregatable A β species, these AD risk factors specify some of the cell surface signal transduction pathways that modulate the phagocytic machinery of microglia.

TYROBP (TYROsine kinase Binding Protein) (also known as DAP12), is a microglial transmembrane signaling polypeptide that contains an immunoreceptor phosphotyrosine-based activation motif (ITAM) in its cytoplasmic domain and is a direct partner/adaptor for immune receptors, including TREM2, CR3, and SIRP\$1 (Signal Regulatory Protein β 1) all of which are independently linked to, or associated with, LOAD [5, 7, 23, 55, 86]. Interaction of TYROBP with its partners forms phagocytosis "active zones" (known as phagocytic synapses) on the surface of microglia. In preparation for phagocytosis, there is a respiratory burst that generates reactive oxygen species (ROS) and appears to involve an interaction between TYROBP and CR3, which in turn interacts with complement component C3 associated with nearby neurites. Mice lacking the complement receptor CR3 or expressing defective TYROBP show reduced ROS production and apoptosis [77]. A recent report demonstrates that the complement pathway can mediate the toxic effects of soluble A β on synapses, and that overactivation of this pathway in AD leads to excessive synapse pruning and early synapse loss [25]. Since the discovery of a link between mutations of *TREM2* and AD, several studies have emerged regarding the role of a loss of function of TREM2 in AD. While these studies have some conflicting results, the most consistent observation is that either *Trem2* deficiency or *Tyrobp* deficiency can cause reduced recruitment of microglial cells around A β plaques. The impact of this reduction in microglia per plaque was interpreted as deleterious in *Trem2* haploinsufficient and *Trem2* deficient mice.

Through a multi-scale integrated computational approach, we and two other independent groups [12, 48, 86] have previously reported TYROBP as a network hub or driver gene in LOAD. Additionally, missense mutations in TYROBP have been recently reported as risk factors for AD [61]. Evidence associating TYROBP to LOAD notwithstanding, it is important to recognize that most TYROBP mutations (as well as TREM2 mutations) represent loss-of-function mutations that result not in AD but in an osteopathy/encephalopathy known as Nasu-Hakola disease (NHD) [59]. One formulation of these data is that the pathogenic mechanism(s) of loss-of-function (nonsense) mutations in TYROBP associated with NHD may cause molecular events that differ from those associated with missense polymorphisms that increase the risk for AD.

Herein, we report the effects of a constitutively null mutation in Tyrobp on the phenotype of an APP/PSEN1 mouse model of AD. In the Tyrobp-null mouse, there is a deletion of exons 3 and 4 resulting in loss of function of the TYROBP protein by deletion of the transmembrane region and part of the cytoplasmic region including the first tyrosine of the ITAM motif [2]. The APP/PSEN1 mouse model [29] expresses APPKM670/671NL/PSEN1^{Δexon9} in neurons and accumulates in the interstitial spaces of the brain fibrillar amyloid that goes on to form typical amyloid plaques accompanied by neuritic dystrophy, age-dependent synaptic loss without neuronal loss, and abnormalities in spatial memory [25, 29, 40, 41]. Since TYROBP expression is increased in the LOAD brain [86], we hypothesized that the APP/PSEN1 phenotype may be improved in the presence of reduced TYROBP levels. Since Tyrobp is not expressed in neurons, our observations in this report describe non-cell autonomous effects wherein signals arising from microglia perturb the homeostasis of nearby neurons or nerve terminals or the pathophysiology of evolving structural intraneuronal or extracellular Alzheimer's pathology.

Methods

Mouse husbandry

The experimental procedures were conducted in accordance with NIH guidelines for animal research and were approved by the Institutional Animal Care and Use Committee (IACUC) at Icahn School of Medicine at Mount Sinai. *APP^{KM670/671NL}/PSEN1^{Δexon9}* (*APP/PSEN1*) and *Tyrobp* knockout (KO) mice were obtained from Jackson Laboratories and Taconic/Merck Laboratory, respectively. *APP/PSEN1* mice were crossed with *Tyrobp* KO mice to obtain *APP/PSEN1* mice heterozygous or KO for *Tyrobp*. Four-month-old male and female mice were killed by decapitation. One hemisphere was collected for immunohistochemical analysis. The second hemisphere was collected for transcriptomic and biochemical analyses.

Immunohistochemical and biochemical analyses

Immunohistochemical and biochemical characterization were performed as previously described [40, 41, 44, 76]. For biochemical analysis, hemibrains were processed via differential detergent solubilization to produce TBS-soluble, Triton-X-soluble, and formic-acid soluble Aβ fractions. For analysis of native oligomeric Aβ peptides, 2 µl protein samples from the TBS-soluble fraction were spotted onto activated/pre-wetted PVDF membrane (0.22 µm; Millipore, Billerica, MA). Membranes were incubated with rabbit pAb A11 (anti-prefibrillar oligomers, 0.5 µg/ml), rabbit pAb OC (anti-fibrillar oligomers and fibrils; 0.25 µg/ml), and mouse mAb Nu-4 (anti-oligomers; $1 \mu g/ml$) [44, 76]. Normalization to total APP/A β signal was achieved by detection of human APP transgene metabolites with the mouse pAb 6E10 antibody (1:1000; Covance, Princeton, NJ). To quantify total Aβ levels, human/rat Aβ 1-40/1-42 ELISA kits (Wako) were used according to the manufacturer's instructions.

For immunohistochemistry, 30 µm thick sagittal sections were incubated with the following antibodies: rabbit anti-Iba1 (1:500; Wako, Richmond, VA), mouse anti-6E10 (1:1000; Covance, Princeton, NJ), and rat anti-CD68 (1:200, mca1957, AbD Serotec BioRad). Sections were then incubated with the appropriate secondary antibody: anti-rabbit Alexa Fluor 488 or Alexa Fluor 568 (1:400; Invitrogen, Carlsbad, CA), anti-mouse Alexa Fluor 568 (1:400; Invitrogen, Carlsbad, CA), and antirat Alexa Fluor 488 (1:400; Invitrogen, Carlsbad, CA) antibodies. ThioflavinS (Sigma-Aldrich, T1892, 1% w/v stock solution) was used for labeling amyloid deposits. For measuring microglia number, Iba1-immunolabeled sections were thresholded and particles analyzed with Fiji (v2.0.0). Sizes of 6E10 immunoreactive plaques and fluorescent intensities were analyzed with Fiji (v2.0.0). The regions of interest were determined by manual tracing. Thioflavin S fluorescence intensity and circularity were analyzed as described [85].

For immunoblotting, membranes were incubated with either anti-CD68 (1:1000, mca1957, AbD Serotec Bio-Rad), anti-phospho-Tau pSer202/Thr205 (1:1000; MN1020, Thermo Fisher Scientific, Waltham, MA), anti-Tau (1:1000; MN1000, Thermo Fisher Scientific, Waltham, MA), anti-Synaptophysin (1:200; ab16659, Abcam, Cambridge, MA), anti-Lamp1 (1:200; ab24170, Abcam, Cambridge, MA), anti-C3 (1:50; ab11862, Abcam, Cambridge, MA), and anti-GAPDH (1:5000; sc32233, Santa Cruz, Dallas, TX) antibodies. Integrated density of immunoreactive bands was measured using MultiGauge Software (FujiFilm). At least two independent western blot analyses were performed and normalized using *APP/PSEN1* female mice as controls.

Behavior analysis

The Barnes Maze test was performed using a standard apparatus [3, 74]. Four-month-old mice were transported from their cage to the center of the platform via a closed starting chamber where they remained for 10 s prior to exploring the maze for 3 min. Mice failing to enter the escape box within 3 min were guided to the escape box by the experimenter, and the latency was recorded as 180 s. Mice were allowed to remain in the escape box for 1 min before the next trial. Two trials per day during 4 consecutive days were performed. The platform and the escape box were wiped with 70% ethanol after each trial to eliminate the use of olfactory cues to locate the target hole. All trials were recorded by video camera and analyzed with ANY-maze video tracking software (Stoelting Co, Wood Dale, USA).

Field electrophysiology

Coronal brain slices containing the hippocampal formation were prepared as previously described [17]. Following anesthesia with isoflurane, brains were rapidly removed and cut into 400 µm thick coronal sections using a vibratome VT1000S (Leica Microsystems, Germany). Brain slices were incubated at room temperature for \geq 3 h in a physiologic ACSF containing 120 mM NaCl, 3.3 mM KCl, 1.2 mM Na₂HPO₄, 26 mM NaHCO₃, 1.3 mM MgSO₄, 1.8 mM CaCl₂, 11 mM Glucose (pH 7.4) and then transferred to a recording chamber perfused with ACSF at a flow rate of ~2 mL/min; experiments were performed at 28.0 ± 0.1 °C. Recordings were acquired with a GeneClamp 500B amplifier (Axon Instruments, Union



«Fig. 1 A decrease in TYROBP protein impairs Aβ deposits compaction, microglial activation, and recruitment around AB deposits in 4-month-old APP/PSEN1 mice. a Images of Iba1-immunostained microglia (green) and 6E10-immunoreactive plaques (red) in frontal cortices and hippocampi of APP/PSEN1, APP/PSEN1; Tyrobp+/and APP/PSEN1; Tyrobp^{-/-} mice. Arrows indicate location of the plaques. Scale bar 500 µm. b Quantification of the number of 6E10-immunoreactive Aβ deposits in cortices and hippocampi (Hip) of APP/PSEN1, APP/PSEN1; Tyrobp^{+/-} and APP/PSEN1; Tyrobp⁻ mice. c Measurements of the size of 6E10-immunoreactive Aß deposits in cortices of APP/PSEN1 mice WT, deficient or null for Tyrobp. d Ouantification of the number of Iba1-immunostained microglia in frontal cortices and hippocampi of APP/PSEN1, APP/PSEN1; Tyrobp^{+/-} and APP/PSEN1; Tyrobp^{-/-} mice. e, f Images of Iba1immunostained microglia (green) and 6E10-immunoreactive plaques (red) and quantification of numbers of cortices plaque-associated microglia located on or within 30 µm radius of 6E10 immunoreactive Aβ plaques in APP/PSEN1, APP/PSEN1; Tyrobp^{+/-} and APP/ *PSEN1; Tyrobp*^{-/-} mice. n = 3-4 mice per group. *Scale bar* 10 µm. g-i Images of thioflavin S-labeled amyloid plaques (g), circularity (h) and quantification of fluorescence intensity (i) of thioflavin S-labeled amyloid plaques from APP/PSEN1, APP/PSEN1; Tyrobp^{+/-} and APP/PSEN1; Tyrobp^{-/-} mice. n = 3-4 mice per group. Scale bar 5 µm. j, k Images of phagocytic microglial marker CD68 (green) and Iba1 (red) co-immunostaining (j) and quantification of fluorescence intensity of CD68 (k) in APP/PSEN1, APP/PSEN1; Tyrobp^{+/-} and APP/PSEN1; Tyrobp^{-/-} mice. n = 3-4 mice per group. Scale bar 30 µm. i Western blot analysis of CD68 in brain protein homogenates from APP/PSEN1, APP/PSEN1; Tyrobp^{+/-} and APP/PSEN1; Tyrobp^{-/-} mice. n = 3-6 mice per group. At least two independent western blot analyses were performed. Representative immunoreactive bands from the same western blot are shown on the right. Oneway ANOVA corrected for multiple comparisons (Tukey) was used for (c, h, i, k) and Two-way ANOVA corrected for multiple comparisons (Tukey) was used for (**b**, **d**, **f**, **l**), *p < 0.05; ***p < 0.001; ****p < 0.0001. Data presented as mean \pm SEM

City, CA) and Digidata 1440A (Molecular Devices, Sunnyvale, CA). All signals were low-pass filtered at 2 kHz and digitized at 10 kHz. For extracellular field recordings, a patch-type pipette was filled with ACSF and placed in the middle third of stratum radiatum in area CA1. Field excitatory postsynaptic potentials (fEPSPs) were evoked by activating Shaffer Collaterals with a Concentric Bipolar Electrode stimulator (FHC, St Bowdoin, ME) placed in the middle third of stratum radiatum 150-200 µm away from the recording pipette. Square-wave current pulses (60 ms pulse width) were delivered through a stimulus isolator (Isoflex, AMPI). Input-output curves were generated by a series of stimuli in 0.1 mA steps. Paired-pulse facilitation was measured by delivering two stimuli at 20, 50, and 100 ms inter-stimulus intervals. Each interstimulus interval was repeated three times and the resulting potentials were averaged. The paired-pulse ratio was calculated by dividing the slope of the second EPSP by the slope of the first EPSP. All results were analyzed by ANOVAs followed by Tukey post hoc tests. Baseline recordings (stable for 20 min) were made every 30 s using stimuli that yielded a response equal to 50% of spike threshold. LTD was induced using a 1-Hz train of 900 bursts, each burst containing three stimuli delivered at 20 Hz, using stimulus strength just superthreshold for evoking a population spike during baseline.

Molecular biological analyses

RNA isolation, library preparation, differential expression analysis and gene set enrichment analyses were performed as described [6, 26, 27, 64, 66, 67].

Computational screen of TYROBP regulating compounds

Drug-induced gene expression fold change was obtained from the Connectivity Map database [42], which consists of 6100 individual experiments, representing 1309 unique compounds. The 6100 individual expression profiles were merged into a single representative signature for the 1309 unique compounds, according to the prototype-ranked list method [28]. Each compound was scored according to the rank of Tyrobp expression fold change within its signature. Compounds were ranked in descending order of Tyrobp expression fold change and used for a secondary enrichment analysis of drug-target associations. For each compound in the drug signature library, referenced drug-target associations [45, 83] and predicted off-targeting [36, 37] were collected. For each of these features, we calculated a running sum enrichment score, reflecting whether that feature was overrepresented among the compounds at the top (associated with Tyrobp upregulation) or at the bottom (associated with Tyrobp down-regulation). Two-tailed p values were based on comparison with 10,000 permuted null scores, generated from randomized drug target sets that contain an equivalent number of compounds to the true set under evaluation, and adjusted using the Benjamini-Hochberg method [6]. Computational screening and chemogenomic enrichment analysis were performed using the R project for statistical computing version 3.2.5 [62].

Data and software availability

Gene expression data generated contributing to the described study will be deposited electronically to the Synapse Web Portal (https://www.synapse.org) in accordance with data sharing policies established by the NIH Accelerating Medicine Partnership (AMP) AD consortium. Specific software will also be made available upon request.

Results

TYROBP deficiency or absence does not modify the number and size of A β plaque depositions nor the number of microglial cells in prefrontal cortex and hippocampus of *APP/PSEN1* mice

We assessed whether TYROBP deficiency or absence modulates A β deposition in *APP/PSEN1* mice. We did not observe differences in number or size of 6E10 immunoreactive plaques in cortices or in the hippocampi of *APP/ PSEN1* mice heterozygous or KO for *Tyrobp* as compared to *APP/PSEN1* mice with normal levels of TYROBP (Fig. 1a–c). It is important to note that 4-month-old *APP/ PSEN1* mice represent an early time point of AD pathology, and all genotypes presented very little A β deposition in the hippocampus as compared to the cortex.

Reduction in total number of microglia has been observed in older TREM2 KO mice with Aβ pathology [30, 81], most likely due to a reduction of microglia proliferation. No differences were observed in younger mice. In our hands, Iba1 immunostaining in 4-month-old *APP/ PSEN1* mice deficient or null for *Tyrobp* did not show differences in the total number of microglia in (pre)frontal cortices (PC) nor in hippocampi as compared to *APP/ PSEN1* mice with normal levels of TYROBP (Fig. 1a, d). Similar results were observed in WT mice with normal or absent TYROBP (see Suppl. Figure 1).

Loss of TYROBP reduces plaque compaction, microglia clustering, and phagocytosis

When 5XFAD mice, which develop rapid and aggressive amyloid pathology and neuronal loss [56], were rendered deficient or null for TYROBP or TREM2, microglial clustering around plaques and plaque compaction were reduced at 4 months of age [85]. We observed decreased microglial recruitment on and around antibody 6E10-immunoreactive A β deposits in the PC of 4-monthold *APP/PSEN1;Tyrobp^{-/-}* mice as compared to *APP/PSEN1* mice with a normal level of TYROBP (Fig. 1e, f). We also observed reduced compaction and fluorescence intensity of thioflavin S reactive plaques (Fig. 1g–i).

We next assessed by immunostaining the level of a phagocytic marker CD68 [88] in the PC of *APP/ PSEN1* mice WT, deficient or KO for TYROBP. In *APP/ PSEN1;Tyrobp^{-/-}* mice, we observed a decreased expression of CD68 in microglial Iba1-positive cells as compared to *APP/PSEN1* mice (Fig. 1j–k). *APP/PSEN1* mice heterozygous for *Tyrobp* did not present a statistically significant reduction of CD68 expression. Accordingly, the level of CD68 in hemibrain protein homogenates was lower in *APP/PSEN1;Tyrobp^{-/-}* as compared with that observed in *APP/PSEN1* mice (Figure i, j). These data support an interpretation that microglial phagocytic activity was reduced in AD mice in the absence of TYROBP (Table 1 for a summary of results).

Aβ levels and oligomeric Aβ in *APP/PSEN1* mice deficient or null for TYROBP

We assessed whether TYROBP deficiency or absence modulates levels of AB species in APP/PSEN1 mice. We measured levels of Aβ40 and Aβ42 in TBS, Triton-X, and formic acid-soluble AB fractions from brains of 4-month-old male and female APP/PSEN1 mice on a Tyrobp heterozygous or null background (Suppl. Figure 2). In males, deletion of one or both *Tyrobp* alleles did not alter levels of A β 40, AB42 or AB42/40 ratio in any of the three fractions as compared to male APP/PSEN1 mice (Suppl. Figure 2a-i). Female *APP/PSEN1;Tyrobp^{-/-}* mice exhibited lower levels of Aβ40 in Triton-X and formic acid fractions relative to APP/PSEN1 mice, resulting in an increase in the AB42/40 ratio in the Triton-X fraction. This was not observed in the formic acid fraction (see Suppl. Figure 2a-i). Notably, female APP/PSEN1 mice WT, heterozygous or knockout (KO) for Tyrobp had higher levels of Aβ40 and 42 in the Triton-X and formic acid fractions when compared to genotype-matched males.

We next assayed oligomeric Aß peptides using antibodies NU-4, A11, and OC antibodies to distinguish among various A_β conformers (Fig. 2a-c). Higher levels of oligomeric Aß reactive with these antibodies have been correlated with impaired cognitive performances in humans and mice [49]. We and others [40, 73] have reported an association of excess levels of NU4-epitope-containing oligomeric A β with deficits in learning behavior in AD mouse models. NU-4 reactive oligomer levels were reduced in both male and female APP/PSEN1 mice with deficiencies in TYROBP as compared to levels observed in APP/PSEN1 mice with normal TYROBP (Fig. 2a). A11 reactive oligomer levels were also reduced in female mice with reduced TYROBP as compared to APP/PSEN1 mice WT for Tyrobp (Fig. 2b). TYROBP level played no obvious role in determining levels of OC epitope-containing oligomeric A β in this system (Fig. 2c). (See Table 1 for a summary of results).

Phospho-TAU, synaptophysin, LAMP1, and C3 levels are altered in *APP/PSEN1* and WT mice with reduced or absent TYROBP

In addition to amyloid deposition, *APP/PSEN1* mice develop hyperphosphorylated microtubule-associated protein TAU (MAPT). We assayed the phosphorylation

Table 1Summary of the assays
performed and results in APP/
PSEN1; Tyrobp^{-/-} vs. APP/
PSEN1, APP/PSEN1; Tyrobp^{+/-}
vs. APP/PSEN1 and APP/
PSEN1; Tyrobp^{-/-} vs. APP/
PSEN1; Tyrobp^{+/-}

Assays	APP/PSEN1; Tyrobp (–/–) vs. APP/PSEN1		APP/PSEN1; Tyrobp (+/-) vs. APP/PSEN1		APP/ PSEN1;Tyrobp (-/-) vs. APP/ PSEN1; Tyrobp (+/-)	
	Females	Males	Females	Males	Females	Males
Αβ40						
TBS fraction	ns		ns		ns	
Triton-X fraction	ns		ns		\downarrow	ns
Formic acid fraction	ns	\downarrow	ns		ns	
Αβ42						
TBS fraction	ns		ns		ns	
Triton-X fraction	ns		ns		ns	
Formic acid fraction	ns		ns		ns	
Αβ42/40						
TBS fraction	ns		ns		ns	
Triton-X fraction	ns		ns		\uparrow	ns
Formic acid fraction	ns		ns		ns	
NU-4	\downarrow	$\downarrow p = 0.08$	ns	\downarrow	ns	
A11	\downarrow	ns	\downarrow	ns	ns	
OC	$\downarrow p = 0.06$	ns	ns		ns	
CD68 (protein)	\downarrow	$\downarrow p = 0.06$	ns		ns	
Phospho-TAU	\downarrow	ns	ns		ns	
Synaptophysin (protein)	ns		ns		ns	
LAMP1 (protein)	\downarrow	\downarrow	↑	ns	\downarrow	\downarrow

 \downarrow Decreased level, \uparrow increased level, *ns* not significant

status of MAPT in WT (nontransgenic) and *APP/PSEN1* mice with normal, reduced, or absent TYROBP (Fig. 2d, e). We observed an apparent increased stoichiometry of TAU phosphorylation in male mice deficient for TYROBP as compared to WT mice (Fig. 2d). Females deficient for TYROBP demonstrated a trend toward increased phosphorylation of TAU as compared to WT mice (p = 0.07). In the presence of mutant *APP/PSEN1* transgenes, there was a reduction in the stoichiometry of TAU phosphorylation in female mice with reduced or absent TYROBP, but no difference in male mice (Fig. 2e).

To examine synaptic integrity, we measured the levels of the presynaptic neuronal marker, synaptophysin (Fig. 2f, g). Synaptophysin was increased in male and female $Tyrobp^{-/-}$ mice as compared to WT mice (Fig. 2f), but no difference was observed between groups in *APP/PSEN1* mice heterozygous-null or homozygous-null for *Tyrobp* (Fig. 2g). Notably, however, LAMP1, a lysosomal protein enriched in dystrophic neurites [15, 21], was decreased in both male and female *APP/PSEN1*; *Tyrobp*^{-/-} mice relative to *APP/PSEN1* alone (Fig. 2h). Excessive activation of the complement system is an early event in AD leading to synapse loss. The level of complement C3 was decreased in female *APP/PSEN1* homozygous-null for *Tyrobp* relative

to those expressing *APP/PSEN1* alone (Fig. 2i). No difference was observed in corresponding male mice (see Table 1 for a summary of results). Despite the relatively minor effect size, likely due to the early stage of AD pathology in 4-month-old *APP/PSEN1* mice, when taken together, these results are consistent with a conclusion that decreased expression of *Tyrobp* may have beneficial effects in the proteinopathy of AD.

Electrophysiological changes in *APP/PSEN1* mice deficient for TYROBP

TYROBP, amyloid, and presenilin proteins play important roles in excitatory synaptic transmission at Shaffer collateral-CA1 pyramidal cell synapses [52, 68]. All of the recombinant mouse models tested in this report showed either altered basal synaptic function, reduced plasticity, or both. Basal synaptic efficiency, as measured by the slope of the input–output relationship, was normal in *APP/ PSEN1* and *APP/PSEN1*; *Tyrobp^{-/-}* as compared to WT mice (Fig. 3a). Interestingly, the slope of the input–output relationship was increased in *Tyrobp^{-/-}* mice compared to WT, *APP/PSEN1*, and *APP/PSEN1*; *Tyrobp^{-/-}* suggesting



Fig. 2 A decrease in TYROBP protein decreases oligomeric Aβ levels and alters phospho-TAU, synaptophysin, LAMP1, and complement C3 levels in 4-month-old *APP/PSEN1* mice. **a**–**c** Hemibrains of male and female *APP/PSEN1* (n = 4–6), *APP/PSEN1; Tyrobp*^{+/-} (n = 3–8) and *APP/PSEN1; Tyrobp*^{-/-} (n = 3–4) mice were processed via differential detergent solubilization to produce fractions of TBS soluble, Triton-X soluble, and formic acid soluble Aβ. Oligomeric Aβ was assessed from the TBS-soluble fraction via dot blot analyses using NU-4 (**a**), A11 (**b**) and OC (**c**) antibodies. **d**–**i** Western blot analysis in brain protein homogenates from 4-month-old male and female mice *WT*, *Tyrobp*^{+/-}, *Tyrobp*^{-/-} and *APP/PSEN1; Tyrobp*^{-/-} mice. **d**, **e** Phospho-tau (AT8 epitope)/total tau ratio for **d** *WT*, *Tyrobp*^{+/-},

Tyrobp^{-/-} mice and **e** *APP/PSEN1*, *APP/PSEN1*; *Tyrobp*^{+/-} and *APP/PSEN1*; *Tyrobp*^{-/-} mice. **f**, **g** Synaptophysin level for **f** *WT*, *Tyrobp*^{+/-}, *Tyrobp*^{-/-} mice and **g** *APP/PSEN1*, *APP/PSEN1*; *Tyrobp*^{+/-} and *APP/PSEN1*; *Tyrobp*^{-/-} mice. **h** Marker of dystrophic neurites (Lamp1) in *APP/PSEN1*, *APP/PSEN1*; *Tyrobp*^{+/-} and *APP/PSEN1*; *Tyrobp*^{+/-} and *APP/PSEN1*; *Tyrobp*^{-/-} mice. **i** Complement C3 in *APP/PSEN1*, *APP/PSEN1*; *Tyrobp*^{+/-} and *APP/PSEN1*; *Tyrobp*^{-/-} mice. At least two independent western blot analyses were performed. Representative immunoreactive bands from the same western blot are shown on the right. *n* = 3–6 mice per group. Two-way ANOVA corrected for multiple comparisons (Tukey) was used for all statistical comparisons in male and female mice, **p* < 0.05; ***p* < 0.01; ****p* < 0.001, *****p* < 0.0001. Data presented as mean ± SEM



Fig. 3 A decrease in TYROBP protein alters excitatory synaptic transmission in the hippocampus in 4-month-old *APP/PSEN1* mice and interacts with the *APP/PSEN1* genotype. In all panels, summary graphs are shown on the *left* and representative traces on the *right*. **a** Basal synaptic function is increased in *Tyrobp^{-/-}* mice, but is unaffected by other transgenic genotypes. The slope of the input/ output relationship was steeper for the *Tyrobp^{-/-}* mice than for all other genotypes (p < 0.05), which did not differ among themselves. **b** *APP/PSEN1* mice showed reduced paired-pulse facilitation (PPF) relative to other genotypes, which did not differ among

an increased basal synaptic activity in absence of TYROBP (Fig. 3a).

We tested the possibly that this effect of *Tyrobp* deletion was presynaptically mediated using paired-pulse

themselves. **c** Synaptically induced long-term depression (LTD) was impaired in all recombinant mice. Analysis over the final 5 min of the recordings showed the most profound deficits for *Tyrobp*^{-/-} and *APP/PSEN1*;*Tyrobp*^{-/-} mice, both of which were significantly less depressed than in *APP/PSEN1* mice. Wild-type (WT) controls showed significantly greater depression than the other genotypes. One-way ANOVA corrected for multiple comparisons (Tukey) was used for statistical comparisons, **p* < 0.05, ***p* < 0.01; ****p* < 0.001. Data presented as mean as mean ± SEM

facilitation (PPF), a short-term form of synaptic plasticity that is sensitive to the probability of transmitter release [9]. PPF was normal in $Tyrobp^{-/-}$ mice (Fig. 3b) suggesting that the increase in basal efficiency observed in

 $Tyrobp^{-/-}$ mice most likely reflects postsynaptic regulation such as increased expression and/or function of synaptic AMPA-type glutamate receptors (AMPARs). These data also raise the possibility of impaired endocytosis of AMPARs [87]. In contrast, PPF was depressed in *APP/PSEN1* mice relative to WT controls, indicating an increase in transmitter release probability. Importantly, *Tyrobp* deletion reversed the deleterious effect of *APP/ PSEN1* on presynaptic function, since PPF was normal in slices from *APP/PSEN1*; *Tyrobp*^{-/-} mice.

We also examined the effects of the different *APP/PSEN1* and *Tyrobp* genotypes on long-term depression (LTD), a persistent form of plasticity whose expression depends on endocytosis of postsynaptic AMPARs [38]. For these experiments, we used a synaptic induction protocol that induces a prominent protein synthesis-dependent "late" phase of LTD [69]. Slices from the *APP/PSEN1* mice showed impaired LTD (Fig. 3c), similar to that reported in older *APP/PSEN1* mice. Similar results were observed following a weaker induction protocol [13] or when late LTD was induced by metabotropic glutamate receptor activation (mGluR-LTD) [84]. LTD was even more impaired in *Tyrobp^{-/-}* and *APP/PSEN1; Tyrobp^{-/-}* mice. Thus, unlike the phenotypes for basal efficiency

and PPF, superimposition of TYROBP deficiency on the *APP/PSEN1* mutations failed to restore normal LTD.

Barnes maze

We next probed the effect of *Tyrobp* deletion on the modulation of spatial learning and memory using the Barnes Maze Test (Fig. 4a–d). The escape latency and distance traveled of *Tyrobp* heterozygous- and homozygous-null mice were identical to WT littermates (Fig. 4a, b). In the presence of *APP/ PSEN1* mutations, deficiency of TYROBP improved learning and memory relative to *APP/PSEN1* with normal TYROBP levels (Fig. 4c, d). This improvement was associated with a reduction in the time spent finding the hidden location (target hole) and a smaller distance traveled in all quadrants. These behavioral data are consistent with a beneficial effect of the *Tyrobp* deletion on the *APP/PSEN1* phenotype.

Differential gene expression analysis of $Tyrobp^{-/-}$ and $Tyrobp^{+/-}$ mice relative to WT mice

Given our extensive database on the regional and diseasestage-specific transcriptomic changes in human LOAD [64, 79, 86], we began by investigating how *Tyrobp* deletion perturbed brain regional transcriptomes. We generated



Fig. 4 A decrease in TYROBP protein improves spatial learning and memory in the Barnes Maze Test in 4-month-old *APP*/*PSEN1* mice. **a-d** 6 groups of 4-month-old mice were used: wild-type (WT) (n = 14), $Tyrobp^{+/-}$ (n = 9), $Tyrobp^{-/-}$ (n = 10), *APP*/*PSEN1* (n = 5), *APP*/*PSEN1*; $Tyrobp^{+/-}$ (n = 9) or *APP*/*PSEN1*; $Tyrobp^{-/-}$ (n = 11). **a, b** Mean latencies to enter the target hole for

a *APP/PSEN1* negative mice and **b** *APP/PSEN1* positive mice. **c**, **d** Mean distances traveled for **c** *APP/PSEN1* negative mice and **d** *APP/PSEN1* positive mice. Two-way ANOVA corrected for multiple comparisons (Tukey) was used for all statistical comparisons, *p < 0.05; **p < 0.01; ***p < 0.001. Data presented as mean \pm SEM

transcriptomic profiles from (pre)frontal cortices (PC) and dentate gyri (DG) (n = 24) for 4-month-old female and male *Tyrobp*^{-/-} and *Tyrobp*^{+/-} mice and compared with WT mice, including sex as a variable. In comparison with non-recombinant (WT) mice, we identified 10 differentially expressed genes (DEG) in the PC, and 28 DEG in the DG of *Tyrobp*^{-/-} mice [false discovery rate (FDR) <0.1] (Fig. 5a). We also identified 2 DEG in the PC and DG of

Tyrobp^{+/-} mice vs. WT mice. *Tyrobp* was the top DEG among the different models and brain areas (logFC = -1.1 and -4.8 in *Tyrobp*^{+/-} and *Tyrobp*^{-/-}, respectively) (Fig. 5b, c). Overall, we observed strong overlap of the DEG between the different brain areas in mice KO for *Tyrobp* (Fig. 5b, c for top 10 DEG in PC and DG, see Suppl. 3 for full DEG results). Thus, eight out of ten DEG in the PC of *Tyrobp*^{-/-} were also differentially expressed

a					d				
Number of DEGs	20 15- 10- 5- 1 0 <i>Tyroby</i> . vs. DG	17 1 1 1 1 1 1 1 1 1 1 1 1 1	8 4 Tyrobp(-+) vs. Tyrobp(++) DG	2 7yrobp(-/-) PC	50 5940- 990 Jo 30 - 1990 Jo 20 - N N 0 A	30 21 PP/PSEN1;Tyrobp(+/-) s. APP/PSEN1	47 37 APP/PSEN1;Tyrobp(-/-) vs. APP/PSEN1 vs	22 10 APP/PSEN1;Tyro , APP/PSEN1;Tyro	Down <u>bp(-/-)</u> <u>bp(+/-)</u>
b	Т	op 10 DEGs in dentate gyri of Tyrot	op(-/-) vs. W	г	е	Top 10 DEGs in A	APP/PSEN1:Tvrobp(+/-) v	s. APP/PSEN1	1
	Symbol Tyrobp Zfp14 Nudt19 Blvrb Kirrel2 Zfp780b Pdcd5 Tdrd12 Gmfg Mrps12	Gene Name TYRO protein tyrosine kinase binding prote ZFP14 zinc finger protein nudix hydrolase 19 biliverdin reductase B kin of IRRE like 2 zinc finger protein 780B programmed cell death 5 tudor domain containing 12 glia maturation factor gamma mitochondrial ribosomal protein S12	logFC -4.7 0.9 -1.2 0.9 1.5 -0.5 -0.5 1.3 1.2 -0.7	FDR 1.86E-24 1.60E-12 2.88E-12 1.32E-10 4.18E-09 7.62E-08 4.86E-06 6.90E-06 1.75E-05 1.84E-05	Symbol Tyrobp Mrps12 Pdcd5 Fzd10 Hipk4 Kctd15 Mc4r Nudt19 Ppp1r14a Prss35	Gene Name TYRO protein tyrosi mitochondrial riboso programmed cell de frizzled class recept homeodomain intera potassium channel t melanocortin 4 rece nudix hydrolase 19 protein phosphatase protease, serine 35	ne kinase binding protein mal protein S12 ath 5 or 10 acting protein kinase 4 etramerization domain conta ptor 9 1 regulatory inhibitor subuni	logFC -1.2 -0.5 -0.5 -1.1 -0.6 1.1 -0.6 1.1 -0.6 it 14A -1.1 0.8	FDR 1.97E-03 3.71E-03 1.29E-03 1.29E-03 1.29E-03 1.29E-03 1.29E-03 1.29E-03 1.29E-03
С	Тс	op 10 DEGs in prefontal cortices of	Tyrobp(-/-) v	s. WT	f	Top 10 DEGs ir	n APP/PSEN1;Tyrobp(-/-)	vs . APP/PSEN	11
	Symbol Tyrobp Nudt19 Zfp14 Pdcd5 Blvrb Mrps12 Hipk4 Sbsn Zfp420 Cep89	Gene Name TYRO protein tyrosine kinase binding protein nudix hydrolase 19 ZFP14 zinc finger protein programmed cell death 5 biliverdin reductase B mitochondrial ribosomal protein S12 homeodomain interacting protein kinase suprabasin zinc finger protein 420 centrosomal protein 89	logFC -4.9 -1.3 0.6 -0.6 0.7 -0.7 -0.7 -0.7 -0.6 -0.5 -0.4	FDR 2.41E-24 1.62E-13 9.50E-08 9.50E-08 1.42E-07 1.48E-04 2.41E-03 5.80E-03 9.84E-03 8.02E-02	Symbol Tyrobp Nudt19 Pdcd5 Mrps12 Gramd1a Bivrb Zfp14 Ppp1r14a Hipk4 Cd33	Gene Name TYRO protein tyrosi nudix hydrolase 19 programmed cell de mitochondrial riboso GRAM domain contr biliverdin reductase ZFP14 zinc finger pr protein phosphatase homeodomain intera CD33 antigen	ne kinase binding protein ath 5 mal protein S12 aining 1A B rotein a 1 regulatory inhibitor subuni acting protein kinase 4	logFC -4.8 -1.5 -0.8 -0.7 0.6 0.9 0.7 it 14A -1.3 -0.7 -1.2	FDR 4.32E-1 8.61E-0 2.03E-0 2.03E-0 1.26E-0 1.54E-0 1.19E-0 2.45E-0 4.56E-0 7.04E-0
					g Top Symbol Tyrobp Nudt19 Map3k14 C230091D Prkdc Blvrb 9430018G Akt2 Acot10 Pak4	10 DEGs in APP/F Gene Name TYRO protein nudix hydrola: mitogen-activ 008Rik RIKEN cDNA protein klinase biliverdin redu 01Rik RIKEN cDNA v-akt murine t acyl-CoA thio p21 protein (C	PSEN1;Tyrobp(-/-) vs. APF tyrosine kinase binding prote se 19 ated protein kinase kinase kin C230091D08 gene , DNA-activated, catalytic po Ictase B 9430018G01 gene hymoma viral oncogene hom ssterase 9 dc42/Rac)-activated kinase 4	P/PSEN1; Tyrob logF ein -3.7 -0.8 nase 14 0.6 -0.4 -0.3 lypeptide -0.3 0.6 -1.0 nolog 2 0.3 -1.0 0.5	(+/-) FDR 4.25E-11 1.12E-02 6.21E-02 1.23E-02 1.23E-02 3.16E-02 3.16E-02 3.23E-02 3.23E-02

Fig. 5 Differential gene expression analysis suggests potential molecular mechanisms associated with TYROBP deficiency. **a**–**c** Differential gene expression analysis in dentate gyrus and prefrontal cortex of *Tyrobp^{-/-}*, *Tyrobp^{+/-}* and *WT* mice. **a** Number of up and down-regulated genes in *Tyrobp^{-/-}* and *Tyrobp^{+/-}* vs. *WT* and *Tyrobp^{-/-}* vs. *Tyrobp^{+/-}*. **b** Top 10 differentially expressed genes in **b** dentate gyrus of *Tyrobp^{-/-}* vs. *WT* and **c** prefrontal cortex of *Tyrobp^{-/-}* vs. *WT*. Bolding highlights differentially expressed genes shared across dentate gyrus and prefontal cortex. RNA sequencing was performed on a total of 47 samples (*Tyrobp^{-/-} n* = 8 males and 8 females, *Tyrobp^{+/-} n* = 8 males and 8 females, and WT *n* = 7–8 samples, eight males and eight females, for each brain regions). All analyses were corrected for sex effect. Differential gene expression threshold was set at fold change ≥1.2 and adjusted *p* value <0.1. *DG*

dentate gyrus, *PC* prefrontal cortex. **d–g** Differential gene expression analysis in prefrontal cortex of *APP/PSEN1;Tyrobp^{-/-}*, *APP/PSEN1; Tyrobp^{+/-}* and *APP/PSEN1* mice at 4-months-old. **d** Number of up- and down-regulated genes in *APP/PSEN1;Tyrobp^{-/-}* and *APP/PSEN1;Tyrobp^{+/-}* vs. *APP/PSEN1* and *APP/PSEN1; Tyrobp^{-/-}* vs. *APP/PSEN1; Tyrobp^{+/-}*. **e** Top 10 differentially expressed genes in *APP/PSEN1; Tyrobp^{+/-}*. **e** Top 10 differentially expressed genes in *APP/PSEN1; Tyrobp^{+/-}*. **v**. *APP/PSEN1; Tyrobp^{-/-}* vs. *APP/PSEN1; Tyrobp^{-/-}* vs. *APP/PSEN1; Tyrobp^{+/-}*. **e** Top 10 differentially expressed genes in *APP/PSEN1; Tyrobp^{+/-}*. **e** Top 10 differentially expressed genes in *APP/PSEN1; Tyrobp^{+/-}*. **e** Top 10 differentially expressed genes in *APP/PSEN1; Tyrobp^{+/-}*. **k** APP/*PSEN1; Tyrobp^{-/-}* vs. *APP/PSEN1; Tyrobp^{+/-}* vs. *APP/PSEN1;*

in the DG of $Tyrobp^{-/-}$. Among them are two genes that have been proposed as early biomarkers of AD: biliverdin reductase B (*Blvrb*) (logFC = 0.9) [54] and Nudix motif 19 (*Nudt19*) (logFC -1.2) [1]. We also noted a strong trend toward down-regulation of *Cd33* in the DG of *Tyrobp*^{-/-} mice (log FC = -0.9, FDR = 0.077). Recent genome-wide studies identified *CD33* as a late-onset AD susceptibility variant [24, 55]. Moreover, CD33 protein is elevated in AD brain and has been associated with amyloid pathology and disease progression [10, 22, 78]. As expected with such small DEG signatures, we did not observe significantly dysregulated Gene Ontology (GO) term enrichments in *Tyrobp* heterozygous- or homozygous-null mice using DAVID, Ingenuity Pathway Analysis (IPA) or gene set enrichment analysis (GSEA).

Differential gene expression and enrichment analysis of *APP/PSEN1;Tyrobp^{-/-}*, *APP/PSEN1;Tyrobp^{+/-}* and *APP/PSEN1* mice

We generated transcriptomic profiles from 23 PC samples from 4-month-old male and female *APP/PSEN1* mice that were either heterozygous- or homozygous-null, or WT for *Tyrobp*. Sex effect was taken into account for all analyses. In comparison to *APP/PSEN1* mice, we identified 84 DEG in *APP/PSEN1;Tyrobp^{-/-}* and 51 DEG in *APP/PSEN1;Tyrobp^{+/-}* (FDR <0.1) (Fig. 5d–g. See Suppl.4 for full DEG results). All of the ten DEG detected in the PC of *Tyrobp^{-/-} vs*. WT mice were also differentially expressed in the PC of *APP/PSEN1; Tyrobp^{-/-}* vs. *APP/PSEN1* comparison. The increased signature size in *Tyrobp^{-/-}* in the *APP/PSEN1* background provides strong independent support for the conclusion that TYROBP is relevant not only in human AD [86] but also in the amyloid-depositing mouse brain AD model.

As above, Tyrobp was the top DEG in both APP/PSEN1 deficient and KO for Tyrobp (logFC = -1.2 and -4.8, respectively). Comparison of APP/PSEN1; Tyrobp^{-/-} vs. APP/PSEN1; Tyrobp^{+/-} highlighted 32 DEG. Interestingly, we found evidence in APP/PSEN1 mice that were heterozygous- or homozygous-null for Tyrobp for several DEG associated with AD and/or memory loss. TYROBP deficiency produced changes in Cd33 expression providing independent confirmation of similar phenomena observed by others using different approaches [12]. Also, Sirt2 expression was increased in APP/PSEN1; Tyrobp^{-/-} $(\log FC = 0.4)$. A SIRT2 polymorphism has been associated with increased LOAD susceptibility [82] and its level of expression is linked with neurodegenerative disease, likely due to its role in lysosome-mediated authophagic turnover [18, 51, 58]. Igfbp2 expression was decreased in APP/PSEN1; Tyrobp^{+/-} (logFC = -0.6). Several proteomic studies aiming to identify AD markers in human sera have reported an increased level of IGFBP2 in AD patients [39, 57]. Moreover, Pedrós et al. have shown an increased expression level of IGFBP2 in hippocampi of an *APP/PSEN1* mouse model similar to that which we used [60]. These data suggest that *Tyrobp*-related modulation of the expression of several AD-related genes only appears in the presence of cerebral amyloidosis and/or *APP/PSEN1* mutations.

To identify biological pathways that may be dysregulated, we performed GSEA using DAVID and IPA (Fig. 6a). Comparisons between APP/PSEN1 KO for Tyrobp and WT for Tyrobp highlighted common themes between DAVID and IPA analyses for perturbation of neurotransmission and ion transport (Fig. 6b and Suppl.5). These included potassium transport, general regulation of transmembrane ion transport, and depolarization and action potential of neurons. Other overlapping themes included neuronal, axonal fasciculation, and synapse assembly. Enrichment analysis with IPA comparing APP/PSEN1; Tyrobp^{+/-} against APP/ PSEN1 detected dysregulation of immune function, including migration, movement, and activation of immune and phagocyte cells (Fig. 6c). Perturbation of metabolic functions was also detected. Using DAVID analysis, we noted a pervasive perturbation of protein phosphorylation signal transduction in APP/PSEN1 mice either heterozygous- or homozygous-null for Tyrobp in comparison to APP/PSEN1 mice on a Tyrobp wild-type background. This is potentially relevant to the role of TYROBP as a phosphotyrosinesignal-based sensor of extracellular debris and instigator and/or organizer of phagocytosis. This is also interesting in light of the evidence that $A\beta$ cerebral amyloidosis in humans and mice is accompanied by hyperphosphorylation of cytoskeletal proteins. Comparisons between APP/ PSEN1; Tyrobp^{-/-} and APP/PSEN1; Tyrobp^{+/-} showed a unique GO term involving stress-activated protein kinase signaling cascade (Fig. 6d). Pathways identified by IPA included production of superoxide, apoptosis, and differentiation/polarization of macrophages. Excessive production of superoxide can induce an uncontrolled oxidative stress leading to increased microglia activation and neuronal apoptosis [80]. Oxidative stress may also promote production and deposition of A β and formation of neurofibrillary tangles [8, 14, 20, 46, 53].

Gene regulatory network analysis

Gene regulatory network analysis is a powerful tool in identifying gene modules pathologically related to complex human diseases including AD [86]. To test if the DE signatures detected in the present study were enriched for any AD networks, we collected the co-expression network modules from our two AD cohorts and overlaid the DEG onto the co-expression network modules. We



Fig. 6 Gene enrichment analysis summary for prefrontal cortex of 4-month-old APP/PSEN1; Tyrobp^{-/-}, APP/PSEN1; Tyrobp^{+/-} and APP/PSEN1 mice at suggests potential molecular mechanisms associated with TYROBP deficiency. a Schematic overview of comparisons between mouse groups. b Enrichment analysis and selected GO terms (DAVID) and diseases and functions (Ingenuity Pathway Anal-

had previously constructed transcriptome-wide gene coexpression networks in different brain regions of postmortem samples from two AD cohorts, the Mount Sinai Brain Bank (MSBB) AD cohort [79] and the Harvard Brain Bank (HBB) AD cohort [86]. Importantly, the age of the mice in this study corresponds to an early stage of AD while our human postmortem co-expression networks from HBB correspond to later disease stages. Although we did not observe enrichments in APP/ *PSEN1*; $Tyrobp^{-/-}$, we found that the DEG down-regulated in PC of APP/PSEN1; Tyrobp^{+/-} (FDR <0.2) were enriched for several sub-networks from both the MSBB and the HBB AD cohorts, including insulin-like growth factor binding, skeletal development, immune system process, anion transport, and particularly the extracellular matrix (see Suppl.6). The up-regulated genes in APP/ *PSEN1;Tyrobp*^{+/-} mice showed enrichment for nucleobase nucleoside and nucleotide metabolic process subnetwork (see Suppl. 6).

Proliferation of antigen presenting cells

ysis) in APP/PSEN1; Tyrobp^{-/-} vs. APP/PSEN1, c in APP/PSEN1; Tyrobp^{+/-} vs. APP/PSEN1, **d** in APP/PSEN1; Tyrobp^{-/-} vs. in APP/ PSEN1; Tyrobp^{+/-}. Enrichments shown were selected for known or suspected relevance to AD pathophysiology. Gene set enrichment threshold was set at p value <0.05. (See Suppl. 5 for full list of enrichments)

Drug repurposing

Through the experiments described above, we identified benefits of TYROBP deficiency on multiple aspects of the phenotype in APP/PSEN1 mice. When those AD model mice were also deficient in TYROBP, beneficial effects in gene expression, phosphorylation of tau, nerve terminal integrity, behavior, and electrophysiology were observed. The data indicate that reduction of *Tyrobp* gene expression could represent a novel computation- and mutation-based, immune-inflammatory therapeutic opportunity for treating or preventing LOAD. Therefore, we probed a comprehensive pharmacopeia database to determine whether safe existing medications would be predicted to reduce levels of Tyrobp mRNA or protein.

To identify small molecule compounds capable of perturbing Tyrobp expression (Fig. 7), we performed a computational screen against a library of drug-induced transcriptional profiles from Connectivity Map [42]. We scored 1309 unique compounds according to the rank of Tyrobp

Fig. 7 Computational analysis of a current pharmacopeia database identified compounds that would be predicted to cause upor down-regulation of TYROBP expression. a Compounds were scored and ranked according to their associated TYROBP expression fold change, and then used as the basis for a secondary enrichment analysis to identify drug targets that associate with up- or down-regulation of TYROBP. Top 10 compounds that \boldsymbol{b} up-regulate and \boldsymbol{c} downregulate TYROBP are shown. Drug targets enriched among compounds that **d** up-regulate, and e down-regulate TYROBP are shown

а		Upregulates TYROBP	6				
			rerences inted its				
			Relo Tary Predict Targets				
			Drug				
		Compound 1					
		Compound 2					
			• •				
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			Identify drug torget enrichmente				
			identity drug target enrichments				
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b							
	Compou	und Dru	ıg class				
	thioguand	osine ANT	IMETABOLITES				
	methapyr	ilene ANT	THISTAMINES FOR SYSTEMIC USE				
	estradiol	EST					
	procainan apigenin	nide AN I	IARRHYTHMICS CLASS I AND II				
	atropine	BEL	LADONNA AND DERIVATIVES, PLAIN				
	minaprine	e ANT	IDEPRESSANTS				
	salsolinol						
С	Salsonitor						
•	Compoi	Ind	Drug class				
	imatinib		OTHER ANTINEOPLASTIC AGENTS				
	benzathin	e benzylpenicillin	BETA-LACTAM ANTIBACTERIALS PENICILLINS				
	diphenylp SC-58125	yraline	ANTIHISTAMINES FOR SYSTEMIC USE				
	lasalocid	-					
	methyldo	oate	ANTIADRENERGIC AGENTS, CENTRALLY	ACTING			
	harpagoside ketotifen		ANTIHISTAMINES FOR SYSTEMIC USE				
	cloxacillin		BETA-LACTAM ANTIBACTERIALS, PENICI	LLINS			
d							
	larget	Name	1	FDR			
		dopamine recep		1.9E-05			
		Botoosium Coloi	pha ZA	0.3E-03			
		rolassium Calci		7.75.05			
		entor Nicotinic Delta Subunit	1.7E-03				
	KYNII Kyoureninase						
	CCR8	CCR8 C-C Motif Chemokine Recentor 8					
	KCNN2	Potassium Calcium-Activated Channel Subfamily N Member 2					
	CHKA	Choline Kinase Alnha					
	ENPEP	olutamyl aminopentidase (aminopentidase A)					
	ADRA1B	adrenoreceptor	alpha 1B	4.9E-02			
	CTSD	Cathepsin D		4.9E-02			
	AOC3	Amine Oxidase.	Copper Containing 3	6.2E-02			
Δ		,					
C	Target		FDR				
	RIPK1	Receptor Interact	ng Serine/Threonine Kinase 1	1.1E-02			
	SELP	Selectin P (granul	e membrane protein 140kDa, antigen CD62)	4.2E-02			

expression fold change (based on comparison to withinbatch vehicle control assays). Top compounds associated with *Tyrobp* up and down regulation are summarized in Fig. 7b.

To explore the pharmacological context of these compounds, we performed a secondary enrichment analysis to identify drug targets that are associated with *Tyrobp* regulation (Fig. 7a, c). We found that compounds that regulate *Tyrobp* expression are enriched for multiple drug targets, including many with known links to Alzheimer's pathology. These include LOAD risk-associated gene Cathepsin D (FDR = 4.9E-02) [16, 70, 72], and *Akt1* (FDR = 6.8E-04), a molecule that is activated in LOAD [65] and is associated with LOAD risk in Chinese Han AD patients with type 2 diabetes [50].

Of the targets enriched among compounds predicted to suppress *Tyrobp* expression (and so potentially representing therapeutic candidates), RIPK1 (FDR = 1.1E-02) was most strongly implicated. Interestingly, RIPK1, a key constituent of the necrosome, was recently shown in an independent study by other investigators to regulate context-dependent regulation of programmed necrosis via formation of an amyloid signaling complex [47]. Experimental validation of these predicted repurposable drugs is underway and will be reported in detail in a future publication.

Discussion

Association of TYROBP with LOAD arose via a multiscale computational network approach [86]. The physical interaction between TREM2 and TYROBP as well as with other LOAD risk factors such as CR3, and SIRP^{β1} [5, 7, 23, 55, 86] provided an important lead for our experimental strategy aiming to validate the important role of TYROBP in the pathogenesis of LOAD. We have previously defined via a multiscale computational network approach TYROBP as a strong candidate for playing the role of a key "hub" or "driver" gene in LOAD [86]. It is worth noting that two independent groups have also identified TYROBP as a driver of LOAD despite having followed different and highly idiosyncratic computational strategies [12, 48]. CD33 is a known AD risk gene and a component of the TYROBP network. The regulation of Cd33 by TYROBP reported herein as well as the regulation of TREM2 by CD33 reported by de Jager and colleagues [12] provide compelling evidence in support of the role of TYROBP as a "driver" gene in LOAD. Capping off the evidence associating TYROBP with LOAD is the recent discovery that missense mutations in the coding region of the TYROBP gene are associated with AD risk [61]. Interestingly, in the same study, in vitro overexpression of the candidate pathogenic p.D50_L51ins14 *TYROBP* variant led to a strong reduction of TREM2 expression [61]. We have previously shown that *TYROBP* expression is elevated in AD brain and mouse models [64, 86], but it was not immediately apparent whether that elevation represented a preexisting, predisposing factor or was a secondary reaction to LOAD pathology. Based on the data presented above, a *Tyrobp* null mutation appears to exert effects that would be characterized as beneficial with respect to both the normal physiology of neurons and the proteinopathy of LOAD.

The effects of Tyrobp knockdown or knockout on AB levels and A β oligomer conformers as defined by epitope content were limited to the reduction of the level of NU-4 and A11 type oligomers in TYROBP deficient APP/PSEN1 mice. There were no consistent statistically significant differences on levels of total AB, AB40, AB42, or on levels of OC type A β oligomers. The relatively minor effect size notwithstanding, it is worth noting that the converging evidence from several laboratories (including our own) is that the NU-4 epitope is the signature of the A β oligomer strain that is most consistently neurotoxic [40, 73]. All and OC oligomer strains are not consistently neurotoxic. As reported above and in one of our previous studies [41], we noted sex differences in A β and oligomer levels suggesting an earlier progression of the disease in female than male APP/PSEN1 mice.

The difference in A β levels observed between the male and female mice is of importance considering the sexual dimorphism observed in the phosphorylation status of TAU in APP/PSEN1 background but not in WT background. Thus, the effect of a decreased Tyrobp expression on the stoichiometry of TAU phosphorylation appeared to be different in the presence or absence of APP/PSEN1 mutations leading to amyloid deposition. Indeed, TYROBP deficiency tends to increase the phosphorylation of TAU on a WT background, but, on a APP/PSEN1 background, loss of TYROBP decreased the phosphorylation status of TAU in female mice in the setting of higher A β loads as compared with males. Although the mechanism(s) by which microglia exert their effects on neuronal tau pathology remains unclear, several reports have linked TREM2 expression and hyperphosphorylated TAU [31, 33, 35]. These reports suggest that TREM2 deficiency could increase tauopathy in human tau-expressing models but could decrease tau pathology in AD mouse models displaying cerebral amyloidosis. Herein we report that a decreased expression of Tyrobp can have beneficial effects on tau pathology and neuronal injury in APP/PSEN1 mouse model of AD. In accordance with our data, Strittmatter and colleagues [75] recently reported that mouse deficient for Progranulin presented an overexpression of Tyrobp network genes correlating with an increased neuronal injury and tau pathology in the absence of amyloid pathology [75].

As mentioned above, no differences were noted in number and size of A^β plaque depositions and the general histological impact of TYROBP deficiency on plaque morphology and microglia recruitment was identical in appearance to that reported by Colonna and colleagues in their studies of TREM2-deficient mice [85]. Indeed, Tyrobp KO mice presented fewer microglia decorating each amyloid plaque without modification in the total number of microglia, and plaques exhibited less compact morphology. However, unlike the Colonna report wherein the reduced numbers of microglia per plaque were predicted to be associated with *increased severity* of the phenotype [85], we observed that this histological appearance was instead associated with beneficial effects on neuritic dystrophy, TAU metabolism, learning behavior, and neuronal electrophysiology. Although beyond the scope of this study, it will be interesting to determine whether overexpression leads to opposite results. In addition, recent papers from Lamb and colleagues [30], Yu and colleagues [32, 34], and Raha-Chowdhury and colleagues [63] raise the possibility that there could be aging-related and/or disease-stage-related changes in the effect of TYROBP. These papers focused on TREM2 and suggest that reduced TREM2 may be beneficial early in life (~4 months) while reduced TREM2 late in life (~8 months) could be detrimental. We are in the process of assessing whether a similar phenomenon occurs with TYROBP.

Electrophysiological assays revealed that the loss of TYROBP normalized some of the synaptic dysfunctions caused by the APP/PSEN1 mutations. The strong increase in basal synaptic efficiency seen in the $Tyrobp^{-/-}$ mice is of particular interest. If observed in isolation, this phenomenon might lead to overactivation of pyramidal neurons and damage, but the same effect could prove protective in the context of LOAD-associated factors that reduce neuronal function. The protective effect of TYROBP deficiency in an early AD context is confirmed by the improvement in the behavioral performance of APP/PSEN1 mice deficient in TYROBP. The effect of the Tyrobp-null background on the electrophysiological findings and gene set enrichment (synapse assembly, ion transport, and neurotransmission) of APP/PSEN1; Tyrobp^{-/-} vs. APP/PSEN1 are in keeping with the growing appreciation for the role of microglia in maintaining normal synaptic physiology [4]. Indeed, in addition to their pro-inflammatory and phagocytic functions, microglia release cytokines including TGF^β and interleukin-1ß that acutely modulate synaptic plasticity at hippocampal synapses [11, 71].

Thus, in a comprehensive panel of transcriptomic, biochemical, electrophysiological, and behavioral paradigms, reduction or ablation of TYROBP prevented the expression of many of the corresponding *APP/PSEN1* phenotypes at 4 months of age. These results would appear to argue against the possibility that early TYROBP deficiency is likely to be a predisposing factor for LOAD. Indeed, these results would indicate that a decrease in TYROBP activity could represent an important therapeutic opportunity either for treating or preventing LOAD or else for slowing or arresting the progression of MCI or early AD to full-blown clinical and pathological LOAD.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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