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2	Transmembrane protein 184B (TMEM184B) promotes expression of synaptic gene
3	networks in the mouse hippocampus
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18	Keywords: TMEM184B, hippocampus, RNAseq, synapse, Alzheimer's Disease

19 Abstract

20 In Alzheimer's Disease (AD) and other dementias, hippocampal synaptic dysfunction 21 and loss contribute to the progression of memory impairment. Recent analysis of human AD 22 transcriptomes has provided a list of gene candidates that may serve as drivers of disease. One 23 such candidate is the membrane protein TMEM184B. To evaluate whether TMEM184B 24 contributes to neurological impairment, we asked whether loss of TMEM184B in mice causes 25 gene expression or behavior alterations, focusing on the hippocampus. Because one major risk 26 factor for AD is age, we compared young adult (5-month-old) and aged (15-month-old) wild type 27 and *Tmem184b*-mutant mice to assess the dual contributions of age and genotype. TMEM184B 28 loss altered expression of pre- and post-synaptic transcripts by 5 months and continued through 29 15 months, specifically affecting genes involved in synapse assembly and neural development. 30 Wnt-activated enhancer elements were enriched among differentially expressed genes, 31 suggesting an intersection with this pathway. Few differences existed between young adult and 32 aged mutants, suggesting that transcriptional effects of TMEM184B loss are relatively constant. 33 To understand how TMEM184B disruption may impact behaviors, we evaluated memory using 34 the novel object recognition test and anxiety using the elevated plus maze. Young adult 35 *Tmem184b*-mutant mice show normal object discrimination, suggesting a lack of memory 36 impairment at this age. However, mutant mice showed decreased anxiety, a phenotype seen in 37 neurodevelopmental disorders. Taken together, our data suggest that TMEM184B is required 38 for proper synaptic gene expression and function but may not be causal for AD and related 39 dementias.

40

41 Introduction

42 Alzheimer's Disease (AD) is a devastating neurodegenerative disease for which we have 43 a paucity of treatment options. In 2022, it is estimated that 10.7% of adults over 65 have 44 Alzheimer's-induced dementia [1]. The burden on patients, their families and health systems is 45 expected to grow significantly in the next 20 years; this trend is already apparent [2]. 46 Alzheimer's Disease causes the accumulation of amyloid beta plagues and 47 neurofibrillary tangles in diverse regions of the brain [3]. Many clinical trials have focused on the 48 depletion of amyloid as a key therapeutic strategy. However, trials in which amyloid plagues are 49 successfully diminished have not produced a reduction or delay in cognitive impairment in trial 50 participants [4]. This highlights the need for the identification of new targets that may contribute 51 to early cognitive impairment and that may offer hope at alternative strategies for stopping the 52 progression of Alzheimer's Disease.

53 Large-scale human genome-wide association and RNAseg analysis to identify network 54 drivers of disease has yielded many plausible candidates that could contribute to AD [5-7]. With 55 these candidates in hand, the task for the AD field now shifts to identifying which candidates 56 cause bona fide alterations in relevant gene expression pathways or in behaviors associated 57 with AD. Through a recent RNAseg analyses, the transmembrane protein TMEM184B was 58 predicted as a candidate protein that could promote AD progression [8]. TMEM184B is a 7-pass 59 transmembrane protein expressed broadly in the nervous system and is thought to play a role in 60 neuronal excitability, synaptic structure, and expression of key developmental and adult 61 pathways involved in neuronal differentiation [9, 10]. Using analysis of the ROSMAP and MAYO 62 clinic human cohorts, TMEM184B was predicted to drive AD-associated gene expression 63 patterns in the dorsolateral prefrontal cortex and temporal cortex, respectively [8]. However, 64 knockdown of TMEM184B in human induced pluripotent stem cell-derived neurons did not 65 directly affect A β levels [5], suggesting that this gene may play a more indirect role.

66 In this study we sought to directly test the hypothesis that TMEM184B regulates gene 67 networks that could contribute to AD, using a mutant mouse model in which the Tmem184b 68 gene is disrupted [11]. We compared how TMEM184B-regulated gene networks are influenced 69 by aging, focusing on the hippocampus as a key location involved in memory. We identified 70 impairments in pathways controlling synapse assembly and function in TMEM184B-deficient 71 hippocampus. We also evaluated memory and anxiety behaviors in these mice. Our results 72 identify a clear effect of TMEM184B on the expression of synaptic gene networks at both young 73 adult and older ages. We also uncovered an unexpected effect of TMEM184B loss on anxiety 74 behavior. However, at young adulthood TMEM184B mutant mice do not have impaired memory. 75 Our work places TMEM184B into a gene regulatory network that could impact synaptic 76 connectivity and influence cognitive function but whose loss likely does not precipitate memory 77 impairment. 78 **Materials and Methods** 79 **Animal Assurances**

All animal experiments are approved by and were performed in accordance with the
 University of Arizona Institutional Animal Care and Use Committee (IACUC) Protocol 17-216.

82 Hippocampal RNA Isolation

83 Male and female mice were humanely euthanized with carbon dioxide, and hippocampi 84 were removed within 10 minutes of euthanasia and immediately frozen for subsequent RNA 85 isolation. Total RNA was extracted using Trizol (Invitrogen) using the manufacturer's protocol. 86 Following initial quality check via Nanodrop analysis, total RNA samples were frozen and 87 shipped to Novogene (Sacramento, CA) for sequencing. Quality control checks were done by 88 Novogene to ensure that sequenced samples had high RNA guality. Sequencing was performed 89 using paired end 150bp reads, 30 million reads/sample, on the Illumina platform. Raw reads 90 data were provided by Novogene.

91 Data Processing and Statistical Analysis

92	Alignment to the mouse genome (mm10.0) was done using Salmon and run on the High
93	Performance Computing cluster at the University of Arizona. Differential gene expression
94	analysis was done using DESeq2 on Galaxy servers (<u>http://usegalaxy.org</u>). PCA plots were
95	generated by Galaxy's DESeq2 default settings, or done in R (https://cran.r-project.org/) using
96	the stats R package.
97	For GO analysis, Panther pathway analysis, and KEGG pathway analysis, ribosomal
98	genes were manually removed from the lists prior to submission. GO analysis
99	(http://geneontology.org/) was performed with a background list of genes consisting of the full
100	list of mapped genes identified by pseudoalignment via the Salmon algorithm from the datasets
101	used for each comparison. KEGG analysis was performed using Enrichr online tools
102	(https://maayanlab.cloud/Enrichr/). Transcription factor binding analysis involved two Enrichr
103	tools: ChEA (2016) and Transcription Factor Perturbations. For both analyses, adjusted P-
104	values were exported and plotted in Graphpad Prism.
105	
106	Data Visualization
107	Fold change values were imported into Cytoscape and mapped onto networks (Biogrid
108	Protein-Protein interaction Network as derived from https://downloads.thebiogrid.org/ or

109 GOCAM Gene Ontology network, as derived from https://github.com/geneontology/gocam-sif-

110 <u>pyexport</u>). To create subnetworks for visualization, we first identified hub proteins that were

111 themselves differentially expressed and also contained multiple differentially expressed genes in

their nearest neighbors. Colorization of levels of fold change used a continuous scale from red

to blue, with grey representing genes not differentially expressed in the comparison groups. For

- biological processes, the default color palette was changed such that darker blues are more
- significant adjusted P-values, while lighter greens to white are less significant.

Graphs were made in either R (ggplot2 package) or in Graphpad Prism. Venn diagrams

117 were created using online tools available at the University of Gent, Belgium 118 (https://bioinformatics.psb.ugent.be/webtools/Venn/). 119 **Novel Object Recognition** 120 Our protocol for novel object recognition was adapted from Leger et al. [12]. Eight to ten 121 mice of each genotype were used for this study; these mice were 5-7 months old and were 122 approximately sex balanced (50/50 or 60/40). Mice were acclimated to the testing room and to 123 the experimenter for two sessions (two days). On day 3, mice were individually placed in an 124 empty square chamber and allowed to explore for 5 minutes. Videos were recorded from above 125 the box in ambient light and with the experimenter outside the room. On day 4 126 (training/association), mice were returned to the chamber to explore two identical objects for ten 127 minutes. On day 5 (24 hours after the day 4 training trial), mice were returned to the chamber 128 which contained one familiar object from the day before, along with one novel object. Ten 129 minutes of object exposure was captured via video. Videos were analyzed offline in Ethovision. 130 To quantify mouse behavior, we used a discrimination index, which is calculated as (total time at 131 novel object - total time at familiar object) / (total object exploration time). Positive values 132 indicate recognition of a novel object.

133

116

134 Elevated Plus Maze

Mice were placed at the distal end of a closed arm and allowed to walk on an elevated X-shaped platform (50cm from the ground, 35 cm arm length, made of gray non-reflective nonodorous poly-methyl methacrylate (PMMA)) for five minutes. Two of the arms of the platform are enclosed by walls, while two are open. The number of times a mouse enters the open or closed arms is recorded. The ratio of open vs. closed entries, and time spent in open versus closed arms, are calculated as an index of anxiety. Benchpads are used under the open arms to cushion any unexpected falls.

TMEM184B expression is required for maintenance of synaptic structure at

142 Results

145

143 **TMEM184B** promotes expression of synapse assembly genes in the mouse

144 hippocampus.

146 neuromuscular junctions in both mouse and fly. We wondered if this role would extend to the 147 hippocampus, a center for learning and memory processing. TMEM184B is expressed in 148 neuronal layers of the hippocampus, suggesting it may contribute here as well. We compared 149 gene expression in the hippocampus in wild type and *Tmem184b* gene-trap mutant mice (which 150 have less than 5% mRNA of Tmem184b remaining [10, 11]) at 5 months of age (Fig 1 and 151 Supplementary File 1). Wild type mice showed high clustering in both principal components (Fig 152 1a), while mutant mice separated well from wild type mice in principal component 1 (PC1). In 153 total, 1153 genes were differentially expressed in *Tmem184b*-mutant hippocampi (Fig 1b-c). 154 When considering those genes with significant hippocampus expression, a group of developmentally important transcripts emerged including Shank1 (involved in post-synaptic 155 156 density scaffolding), Neurexin 1 (involved in synaptic assembly and adhesion), and 157 Somatostatin (a marker of subtypes of interneurons). 158 To evaluate the biological processes likely affected by TMEM184B in the hippocampus, 159 we used gene ontology analysis to pinpoint processes with significant enrichment of 160 differentially expressed genes (DEGs) in our dataset (Fig 1d). This showed a large predicted 161 effect on synapse assembly and neural development. Other processes that may be affected in 162 Tmem184b-mutant mice include memory/cognition, synapse function, behavior, and neuronal 163 cell death. We performed pathway analysis on these data by guerying the KEGG database (Fig 164 1e). This analysis identified many key pathways affecting both glutamatergic (excitatory) and 165 GABAergic (inhibitory) synapses. Furthermore, it suggested that the pathways dysregulated in 166 *Tmem184b*-mutants were enriched for those contributing to neurodegenerative diseases 167 including Parkinson's disease and Alzheimer's Disease. Taken together, our analysis of 7

transcriptomic changes in *Tmem184b*-mutant hippocampus predicts significant dysregulation of
synapses that may be linked to neurological disease.

170

171 Synaptic Protein-Protein Interaction Networks and Biological Processes rely on

172 **TMEM184B** for proper hippocampal expression.

173 We sought to take a closer look at the relatedness of genes that were differentially 174 expressed in the pathways described above, specifically synapse assembly and neural 175 development. Using Cytoscape, we visualized our gene expression changes on the predicted 176 mouse protein-protein interaction network annotated by Biogrid (Fig 2-B). We noticed a 177 significant enrichment of altered transcripts of proteins that are predicted to interact with two key 178 neurodevelopmental proteins: Neurogenin3 and with Shank1. However, contrary to our 179 expectations, Tmem184b-mutant mice show upregulation of proteins in these networks (shown 180 in circles with pink and red shading). Neurogenin 3 is a transcription factor that promotes 181 dendritogenesis in cultured hippocampal neurons [13]. Shank1, along with its family members, 182 are critical scaffolds for postsynaptic density proteins and glutamate receptors, and they are 183 implicated in autism spectrum disorders [14, 15]. The upregulation of these proteins could be a 184 direct effect of TMEM184B or a compensatory effect following other synaptic disruptions; our 185 data does not distinguish between these possibilities. Nevertheless, the correct levels of these 186 factors are crucial for maintenance of proper circuit assembly and function; both up- and down-187 regulation of both proteins is deleterious [13, 16].

In addition to mapping our data onto known protein-protein interactions, we also mapped our data onto gene ontology networks and assessed which neuronal or synapse-related processes were predicted to be altered by TMEM184B expression. The most differentially expressed GO Biological Processes associated with synapses were in synaptic transmission and plasticity. This prediction is consistent with data showing that TMEM184B mutation causes hyperexcitability at glutamatergic synapses in *Drosophila* [9]. Within the search term "neuron"

194	these processes were enriched for neurogenesis and projection (axon/dendrite) development.
195	Overall, our analysis suggests that TMEM184B maintains appropriate expression of
196	interconnected networks of synaptic and neurodevelopmental genes in the hippocampus.

197

198 Analysis of Transcription Factors Upstream of Hippocampal Differentially Expressed

199 Genes implicates Wnt signaling alterations.

200 To understand why loss of TMEM184B causes gene expression changes in the 201 hippocampus at 5 months, we performed an *in silico* analysis to identify enriched transcription 202 factor target genes in our DEGs using two platforms. In the first (ChEA 2016), we identified four 203 transcription factors with enriched targets (DMRT1, RARB, ZFP281, and TCF7) (Fig 3A-B). 204 DMRT1 is not expressed in the brain and so is unlikely to have an effect. ZFP281 is a 205 transcriptional repressor that maintains the pluripotency of stem cell populations and is highly 206 expressed. RARB, or retinoic acid receptor beta, is a steroid hormone receptor responsive to 207 retinoic acid (RA) and plays roles in embryonic development. Disruptions to RARB lead to 208 neurodevelopmental delay and are associated with autism spectrum disorder [17]. TCF7 is a 209 canonical Wht signaling pathway mediator [18]. In a separate analysis, we used the EnrichR 210 platform to examine the overlap between DEGs from our data set with mouse transcription 211 factor manipulations performed by others. We identified many transcription factors whose gene 212 expression alterations significantly resemble that of *Tmem184b*-mutant hippocampus. 213 Interestingly, multiple associations were identified with transcription factors known to promote 214 Wnt signaling (NEUROD1, TCF3, PLAGL2) [19][20, 21]. This is consistent with prior work 215 suggesting that Wnt pathway dysfunction could participate in TMEM184B-dependent 216 phenotypes [10]. Wnt/β-catenin signaling is critical for hippocampal neurogenesis [22, 23], so 217 this suggests that one effect of *Tmem184b* disruption may be loss of neurons. Other factors, 218 such as NFIA, are known contributors to stem cell maintenance and proliferation. Adult 219 neurogenesis in the hippocampus is required for long-term spatial memory in mice [24]. Taken

together, this analysis suggests a possible role for TMEM184B in the generation of hippocampalneurons via promotion of Wnt pathway signaling.

222

223 Identification of Common and Unique Aging Signatures in *Tmem184b*-mutant

224 hippocampus

225 The biggest risk factor for the development of Alzheimer's Disease is advanced age. To 226 identify unique signatures of TMEM184B that may affect the aging brain, we compared genes 227 that were altered upon aging (15 months of age) in wild type mice with those in *Tmem184b*-228 mutant mice. The *Tmem184b* transcript itself was not significantly altered across wild type 229 aging, although it was slightly reduced in older mice ($log_{2}FC = -0.13$; Adjusted P value = 0.074). 230 After filtering out ribosomal transcripts, a total of 2171 genes were significantly altered in 231 15-month-old Tmem184b-mutant mice compared to age-matched wild types (Supplementary 232 File 2). Of these, 222 transcripts were altered at both ages (Fig 4A-B and Supplementary File 233 3). We identified the most enriched processes in this group using gene ontology (Fig 4C-D). 234 This analysis confirmed the strong effects of TMEM184B loss on synaptic processes, including 235 postsynaptic density organization, glutamate receptor and neuroligin binding activity, and 236 synaptic plasticity. Interestingly, both young and aged TMEM184B mutant mice have a small but 237 significant upregulation of ApoE transcripts (log₂FC of 0.23 and 0.30, respectively). This 238 comparison confirms that, throughout aging, TMEM184B continues to influence synaptic gene 239 expression.

If we compare 5-month-old and 15-month-old mutant mice, very few genes are significantly different (104 total) (Fig 4E and Supplementary File 4). In this group, no biological pathways, molecular functions, or cellular components reached statistical significance (all had adjusted p values > 0.05). This indicates that the changes that have occurred in TMEM184B mutant mice primarily occur by 5 months, and gene expression is relatively constant beyond that point. We used the Molecular Signatures database (MSigDB) to query whether alterations

in this small group of genes were similar to other published transcriptomic datasets. This
analysis revealed that the most similar datasets are mouse models of neurological disease,
including two Huntington's Disease models (R6/2 and Q175) and spinal cerebellar ataxia
models caused by disruption of Ataxin-1 (Fig 4F). This suggests that the few genes that are
altered by aging in TMEM184B mutants may affect similar pathways to those affected in HD and
Ataxia.

252

253 Somatostatin, and other neuronal genes, are similarly regulated in both central and

254 peripheral neurons

255 To evaluate common signatures of TMEM184B-dependent gene expression across 256 disparate types of neurons, we compared the differentially expressed genes (adjusted p < 0.05) 257 between our 5 month hippocampus data set and one we have previously reported from 6 month 258 adult dorsal root ganglia [10]. Among differentially expressed genes, we identified 12 genes 259 (including *Tmem184b* itself) that were similarly regulated (Fig 5 and Supplementary File 5). In 260 many cases, the fold change in the hippocampus was somewhat less than that in the DRG, 261 which could reflect a greater percentage of non-neuronal cells in the hippocampus that may 262 dilute the overall fold change. Nevertheless, we noticed that Somatostatin, a neuropeptide used 263 as a marker of GABAergic inhibitory interneurons in the hippocampus (but which marks 264 excitatory, pruriceptive neurons in the DRG), was downregulated in both data sets, suggesting a 265 common means of regulation in these two tissues. One common upregulated gene, Tsukushi 266 (*Tsku*), is a negative regulator of the Wnt pathway [25]. Taken together with our earlier 267 analyses, this again implicates a loss of Wnt signaling in TMEM184B-associated phenotypes. 268 Finally, Bc1, a long non-coding RNA that is upregulated in TMEM184B-deficient neurons, has 269 been shown to promote translation of amyloid precursor protein (APP) and cause spatial 270 learning and memory defects [26].

271

272 TMEM184B loss does not affect spatial memory but alters anxiety behaviors in middle

273 aged mice

274 Finally, we sought to evaluate how these gene expression changes in the hippocampus 275 affect behaviors driven by this brain region, including anxiety and object-oriented memory. In 276 prior work, we established that six-month-old *Tmem184b*-mutant mice show no deficiencies on 277 the rotarod assay, but show difficulty in an inverted screen test, indicating that some 278 sensorimotor impairment is present [10, 11]. Prior to performing memory assays, we first 279 evaluated mobility of mice in an open field, using the same arena we planned to use for spatial 280 memory testing. We did not observe any significant differences in mobility (total distance 281 traveled) (Fig 6A), indicating that we could use this paradigm and arena for novel object 282 recognition, a classical object-oriented memory paradigm. However, we did not see any 283 difference between wild type and *Tmem184b*-mutant mice in their ability to recognize novel 284 objects (both showed a positive discrimination index, indicating more time at the novel object) 285 (Fig 6B). This indicates that at six months of age, TMEM184B loss does not alter memory. 286 Next, we evaluated anxiety using the elevated plus maze. Anxious mice spend more 287 time in closed arms (with walls) than in open arms (without walls). Surprisingly, we identified a 288 decreased anxiety overall in *Tmem184b*-mutant mice when compared to wild types. 289 *Tmem184b*-mutant mice spent considerably more time in the open arms of the maze (Fig 6C). 290 This indicates that TMEM184B promotes appropriate anxiety levels in the elevated plus maze 291 assay.

292

293 Discussion

TMEM184B was predicted in human transcriptome analyses to be associated with deleterious gene expression that could drive the progression of Alzheimer's dementia [8]. Our experiments sought to evaluate this hypothesis using RNAseq analysis across aging in young

297 and aged *Tmem184b*-mutant and wild type mice. Our results show that TMEM184B has 298 significant effects on synaptic gene expression in the hippocampus in both young and old mice. 299 Alteration of post-synaptic density gene transcripts occurred in all of our pathway 300 analyses. Proteins in this group, such as SHANK1, SHANK3, and PSD95/DLG4 contribute to 301 the scaffolding of neurotransmitter receptors. Interestingly, transcripts of all three of these genes 302 are over-expressed in *Tmem184b*-mutant mice. This upregulation of mRNA could reflect a 303 compensation for presynaptic dysfunction, an inappropriate dendritic overgrowth, or a failure to 304 prune exuberant synapses. In fly models of mutation of the TMEM184B ortholog, *Tmep*, extra 305 active zones and synaptic release sites (boutons) are observed at neuromuscular junctions [9], 306 suggesting that additional synapses may be a common effect of TMEM184B disruption. Further 307 detailed analysis of hippocampal synaptic morphology and physiology will be necessary to 308 parse apart these possibilities.

309 Somatostatin (Sst), a common downregulated gene across both central and peripheral 310 TMEM184B-deficient data sets, plays many distinct roles in the nervous system. In dorsal root 311 ganglia, it is a neurotransmitter for a subset of itch-activated sensory neurons [27]. TMEM184B 312 loss in DRG neurons causes disruption of sensation of interleukin-31, a key cytokine involved in 313 atopic dermatitis (eczema) that is detected exclusively by SST+ neurons [27, 28]. In contrast, in 314 the dentate gyrus, somatostatin reduction occurs in genetic models of temporal lobe epilepsy 315 [29]. Reintroduction of SST in this model reverses the epileptic phenotypes, arguing for a causal 316 role in suppression of seizures. Drosophila models of TMEM184B deficiencies show ectopic 317 firing and elevated synaptic calcium, indicating a role in restraining excitability [9]. Thus, 318 phenotypes of SST and TMEM184B overlap substantially; our determination that TMEM184B 319 regulates Sst expression in multiple cell types could explain this concordance. It would be of 320 future interest to determine whether restoration of SST in TMEM184B mutant mice could restore 321 normal gene expression and/or behaviors.

322 Another common theme among our data is the connection between TMEM184B and 323 What pathway regulation. In our prior work, we showed that TMEM184B expression positively 324 influences the expression of many Wnt signaling components, as well as their downstream 325 targets, in developing somatosensory neurons [10]. In this work, we also find dysregulation of 326 Wnt components in the hippocampus, including the loss of Wnt pathway targets as well as 327 upregulation of Wnt pathway inhibitors such as Tsukushi. Wnt signaling deficiency contributes to 328 AD phenotypes in mouse models [30], and it has also been identified as a key pathway altered 329 in human AD samples [31]. While our data does not directly link TMEM184B to the development 330 of AD pathology, it suggests that TMEM184B dysfunction disrupts synaptic gene regulatory 331 networks and also influences expression of genes with more direct links to AD. 332 The mechanism of action of the TMEM184B protein is yet unknown. Clues to its role 333 include localization to endosomes and synaptic vesicles, accumulation of multilamellar 334 structures in mutant presynaptic terminals, and additional autophagosomes and lysosomes in 335 skeletal muscle in its absence. Because Wnt signaling occurs within endosomes and

contributes to dendritic growth, we hypothesize that a failure of Wnt signaling (perhaps due to ablockade in the endolysosomal pathway) may account for some of the phenotypes we observe.

While we did not complete an exhaustive behavioral analysis, we found that TMEM184B disruption causes reduced anxiety in the elevated plus maze paradigm. Reduced anxiety is seen in some models of neuroatypical development including autism spectrum disorders and in mice predisposed to depression [32, 33]. We speculate that TMEM184B loss may disrupt anxiety circuits to cause this phenotype.

We did not observe memory impairment when TMEM184B was disrupted in 6-month-old mice. It is still possible that older mice may show impairments. However, perhaps more likely, we suspect that TMEM184B disruption itself is not causative for memory loss. Instead, we hypothesize that compromising TMEM184B function, in the presence of other susceptibility loci, may increase the likelihood of developing dementia. Given its strong effects on synaptic gene

348	expression networks as well as the disruptions to synaptic morphology and function seen in
349	other studies, it will be of future interest to directly evaluate synaptic function via electrical
350	recording in the hippocampus and to evaluate its role in the context of mouse models of AD.
351	
352	Data Availability
353	The data sets generated and analyzed in the current study have been deposited to NCBI
354	GEO at the following accession: GSE204831. All custom-written code in R for RNAseq analysis
355	is available at GitHub (https://github.com/eriklarsen4/ggplot-scripts.git and
356	https://github.com/martharcb/). Dorsal root ganglion data to which we compared our
357	hippocampal RNAseq data can be found on NCBI GEO (GSE154316).
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370	Conflict of Interest
371	The authors do not report any conflicts of interest pertaining to the content of this manuscript.
372	
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472 **Figure Legends**:

473

474	Figure 1. TMEM184B-dependent gene expression alterations in the mouse hippocampus.
475	A, Principal component analysis (PCA) of wild type and mutant samples taken at 5 months of
476	age shows strong clustering of wild types (black circles), and loose clustering of mutant samples
477	(green circles). B , volcano plot showing both up- and down-regulation of genes based on
478	differential expression analysis. Purple indicates adjusted P-values of $p < 0.05$. C , MA-plot of
479	base mean expression value versus Log_2 (Fold Change) from DeSeq2. Red, Adjusted p < 0.05;
480	blue, adujsted p < 0.01. Selected genes of interest are labeled. D , GO Biological Process Fold
481	Enrichment (FE) scores of categories above 2.0 FE. Numbers indicate average FE;
482	parentheses denote number of processes in that category. E , KEGG Pathways analysis (done
483	using Enrichr). Top 10 pathways are shown, ranked by p-value.
484	
485	Figure 2. TMEM184B loss disrupts key synaptic gene networks. A-B, mouse protein
486	interaction networks as defined by BioGRID. Node color represents log_2 of fold change (darker
487	red indicates higher over-expression in Tmem184b-mutant; blue shades indicate
488	downregulation in <i>Tmem184b</i> -mutant). Circles indicate adjusted P-values where p < 0.05;
489	rectangles indicate p > 0.05. Edge lengths have no significance. C-D , GO Biological Process
490	network maps (from GoCAM) showing terms (gene sets) having significant alteration in
491	Tmem184b-mutant mice versus controls. Darker blue indicates stronger fold change, while
492	larger nodes indicate stronger significance (all nodes shown have Adjusted p < 0.05). Edge
493	lengths have no significance. GO Biological Process terms disrupted in Tmem184b-mutant
494	hippocampi identified with the search term "synapse" (in C) or "neuron" (in D).
495	
496	Figure 3. Analysis of Transcription Factors Upstream of Hippocampal DEGs identifies

497 Wnt pathway alterations. A, Chromatin Enrichment Analysis (ChEA) for mouse transcription

factors (TFs) with binding in promoters of differentially expressed genes from *Tmem184b*mutant hippocampi at 5 months of age. Shown are TFs for which enrichment scores were
statistically significant (Adjusted P < 0.05). **B**, Base mean expression in 5-month-old
hippocampus of each transcription factor in A, graphed against its adjusted P-value. **C**, top 10
mouse data sets from transcription-factor-manipulated genetic backgrounds that show
significant overlap with TMEM184B differentially expressed genes. Adjusted P values calculated
by Enrichr.

505

506 Figure 4. Gene Expression Changes in 15-month Tmem184b-mutant vs wild type mice. A, 507 MA-plot of gene expression changes between 15-month-old *Tmem184b*-mutant and wild type 508 hippocampus. Red dots show genes with adjusted p < 0.05, while blue dots show adjusted p < 0.05509 0.01. B, Venn diagram showing overlap between 5-month and 15-month genotype-induced 510 changes. **C-D**, evaluation of enrichment among the 222 overlapping genes using gene ontology 511 for Biological Processes (C) and Cellular Components (D). Shown is Combined Score reported 512 by Enrichr. This score takes into account both P-value and Odds Ratio. Asterisks indicate level 513 of statistical significance (Adjusted P-value, **** <0.0001, *** < 0.001, ** < 0.01). **E**. Four-way 514 comparison of differentially expressed genes (no ribosomal gene filtering). F, gene set 515 categories identified by Molecular Signatures Database (MSigDB) among genes different 516 between 5- and 15-month Tmem184b-mutant mice. Abbreviations: HD, Huntingon's Disease 517 model mice; Oligo, oligodendrocytes; Ataxin, Ataxin-1 mutant mice. The full list of molecular 518 signatures is shown in Supplementary Table 5.

519

520 Figure 5. TMEM184B similarly regulates somatostatin and other transcripts in both

521 central and peripheral neurons. Data for DRG is available at NCBI GEO (NCBI GEO

522 GSM4668859) Log₂FC of genes identified in each data set (hippocampus, black bars; adult

523 DRG, grey bars). A full list of genes differentially regulated in both data sets can be found in524 Supplementary Table 6.

525

526	Figure 6. Memory and Anxiety Behaviors in Tmem184b-mutant mice. All graphs show mice
527	of both sexes and are approximately sex balanced. A, Open field habituation over five minutes
528	for mice 5-7 months of age. Total distance traveled in centimeters. WT, wild type, Mut,
529	Tmem184b-gene trap mutant mice. B, Memory test at 24 hours using the novel object
530	recognition task. Discrimination index is calculated as described in materials and methods. Each
531	dot shows an individual mouse. N = 10 wild type and 8 mutant mice. C , measurement of anxiety
532	using the elevated plus maze. For all graphs, statistical evaluation used unpaired t-test with
533	Welch's correction. Numbers in bars (in A and C) indicate sample size. Error bars show
534	standard error of the mean (SEM) for all panels.
535	
536	
537	Supplementary Material
538	
539	Supplementary File F1. DESeq2 output of differential gene expression analysis between the
540	hippocampi of 5-month-old Tmem184b-mutant mice and age-matched wild types.
541	
542	Supplementary File F2. DESeq2 output of differential gene expression analysis between the
543	hippocampi of 15-month-old Tmem184b-mutant mice and age-matched wild types.
544	
545	Supplementary File F3. Differentially expressed genes in Tmem184b-mutant mice common to
546	both the 5-month-old and 15-month-old comparison groups. Ribosomal transcripts have been
547	omitted.
548	

549	Supplementary File F4. DESeq2 output of differential gene expression analysis between the
550	hippocampi of 15-month-old Tmem184b-mutant mice and 5-month-old Tmem184b-mutant mice.
551	
552	Supplementary File F5. Analysis of 104 differentially expressed genes across aging in
553	Tmem184b-mutant mice using the Molecular Signatures Database (MsigDB).
554	
555	Supplementary File F6. All genes identified as differentially expressed in both adult dorsal root
556	ganglia and hippocampus of Tmem184b-mutant mice compared to wild type mice.
557	
558	



-log₁₀(P-Val)







Shank1 Subnetwork



GO Biological Process **D** "Synapse" Subnetwork

regulation of

synaptic transmission

synaptic

plasticity

regulation of transmission of

regulation of

bi

nerve impulse

regulation of

transmission of

nerve impulse

synaptic

transmission

cell-cell

signaling

GO Biological Process "Neuron" Subnetwork



С



Α

С

Ε





F





