Triggering receptor expressed on myeloid cells 2 (TREM2) is an orphan immune receptor expressed on cells of myeloid lineage, such as macrophages and microglia. The rare variant R47H of TREM2 is associated with an increased risk for Alzheimer’s disease, supporting the hypothesis that TREM2 loss of function may exacerbate disease progression. However, a complete knockout of the TREM2 gene in different genetic models of neurodegenerative diseases has been reported to result in both protective and deleterious effects on disease-related end points and myeloid cell function. Here, we describe a Trem2R47H transgenic mouse model and report that even in the absence of additional genetic perturbations, this variant clearly confers a loss of function on myeloid cells. The Trem2R47H variant—containing myeloid cells exhibited subtle defects in survival and migration and displayed an unexpected dysregulation of cytokine responses in a lipopolysaccharide challenge environment. These subtle phenotypic defects with a gradation in severity across genotypes were confirmed in whole-genome RNA-Seq analyses of WT, Trem2−/−, and Trem2R47H myeloid cells under challenge conditions. Of note, TREM2-activating antibodies that boost proximal signaling abrogated survival defects conferred by the variant and also modulated migration and cytokine responses in an antibody-, ligand-, and challenge-dependent manner. In some instances, these antibodies also boosted WT myeloid cell function. Our studies provide a first glimpse into the boost in myeloid cell function that can be achieved by pharmacological modulation of TREM2 activity that can potentially be ameliorative in neurodegenerative diseases such as Alzheimer’s disease.

Triggering receptor expressed on myeloid cells (TREM2) is expressed on cells of myeloid lineage, including macrophages and microglia (1, 2). TREM2 is an orphan immune receptor with a short intracellular domain and functions by signaling through its adaptor partner, DAP12 (3). Mutations in both TREM2 and DAP12 have been linked to the autosomal recessive disorder Nasu–Hakola disease, which is characterized by bone cysts, muscle wasting, and demyelination (2). More recently, variants in the TREM2 gene have been linked to increased risk for Alzheimer’s disease (AD) (4, 5) and other forms of dementia, including frontotemporal dementia (4–6). In particular, the R47H variant has been identified in genome-wide studies as being associated with increased risk for late-onset AD with an overall adjusted odds ratio (for populations of all ages) of 2.3, second only to the strong genetic association of APOE with Alzheimer’s. The R47H mutation resides on the extracellular Ig V-set domain of the TREM2 protein and has been shown to impact lipid binding and uptake of Aβ (7–10), suggestive of a loss of function linked to disease. Two recent publications have demonstrated defects in the autophagy/lysosomal pathway (11) and decreased capacity of R47H microglia to compact plaques (12). In general, however, the effect of the variant on TREM2 expression and myeloid cell functioning and the ability to modulate the gene to have a beneficial effect in disease are emerging areas of TREM2 biology. Most preclinical models that have been designed to study TREM2’s role in modulating myeloid cell function and disease progression in AD have employed complete gene knockouts that are probably not representative of a physiologically relevant state. Different knock-out mice crossed with traditional AD mouse models (APP/PS1 and 5XFAD) have resulted in conflicting data with respect to the role of TREM2 in Alzheimer’s disease; Wang et al. (8) have proposed a loss of function for the gene in disease based on decreased microglia survival and reduced ability to cluster around Aβ plaques and clear them (APP/PS1 and 5XFAD crossed with Colonna knockouts). Jay et al. (13) have provided evidence supportive of an amelioration of the neurodegenera-
tive phenotype upon Trem2 deletion in their models, including an increase in phagocytosis and anti-inflammatory cytokines and an overall decrease in amyloid burden (APP/PS1 crossed with NIH KOMP knockouts, hereafter referred to as Trem2R47H). The effects were subsequently shown to be potentially temporally regulated based on disease progression (14). Similarly, conflicting results have been reported in tauopathy models also with an exacerbation in disease end points in one model (15) and an improvement in another (16) upon gene deletion. At the time of submission of this manuscript, Song et al. (17), described for the first time a deleterious effect of human R47H TREM2 when introduced into 5×FAD mice, noting subtle cell intrinsic and extrinsic TREM2-dependent effects. The authors noted differences in plaque load and microglial function between KO-5×FAD mice and human R47H TREM2–5×FAD mice. A more comprehensive understanding of the effect of the variant on myeloid cell function in general, especially in the absence of additional genetic perturbations that are used to generate neurodegenerative disease models, may help clarify some of the contradictory data pertaining to the role of the protein in progressive, neurodegenerative diseases. We describe below an R47H transgenic model and comprehensively characterize the loss-of-myeloid cell function conferred by the R47H variant. We also demonstrate that antibodies that activate proximal TREM2 signaling rescue some of these cumulative defects and even boost WT TREM2 activity. Our studies provide the first glimpse into pharmacological modulation of TREM2 as a means to modulate myeloid cell function in neurodegenerative diseases.

Results

Gene-edited Trem2R47H and Trem2–/– mice are specifically modified in the TREM2 gene with no off-target effects on other TREM genes in the locus

To address some of these fundamental questions linked to TREM2 biology and the effect of the R47H variant on TREM2 functioning, we have generated Trem2–/– and Trem2R47H knockin mice. The TREM gene locus in humans and mice includes multiple TREM and TREM-like genes with both ITAM- and ITIM-associated functions (18) (Fig. S1A). Regulatory elements located within the TREM2 gene can impact the expression of other TREM genes. To specifically target the TREM2 gene without perturbing additional regulatory elements, we used a gene editing–based approach to generate Trem2–/– or Trem2R47H mice. The Trem2–/– strain was generated by engineering a deletion in exon 1 of the Trem2 gene, and the Trem2R47H1 strain was generated by engineering a point mutation at residue 47 in the mouse Trem2 gene (Fig. S1, B and C), analogous to the human variant. Detailed qPCR analyses of brain homogenates from the gene-edited Trem2–/–, Trem2R47H, and commercial NIH KOMP mice confirmed a comparable loss of the gene in the knockouts and Trem2 expression comparable with WT age-matched controls for the Trem2R47H mice (Fig. 1, A–C). We further confirmed that upon LPS stimulation, R47H Trem2 was down-regulated in mouse brains comparable with age-matched WT controls, whereas no detectable levels of Trem2 mRNA were observed in the Trem2–/– animals under any of the conditions tested (Fig. 1, A–C). Similar and expected trends in expression patterns were observed in WT, gene-edited Trem2–/–, and Trem2R47H mice with respect to other Trem genes in the locus, including Trem1 (Fig. 1, D and E), Trem1l (Fig. 1, G and H), and Trem2l (Fig. 1, J and K) under basal and LPS-stimulated conditions. Of note, we and others (19) have observed unexpected changes in other Trem genes in the Trem locus under basal and LPS-stimulated conditions in the Trem2R47H1–/– mice; this includes a less significant Trem1 response compared with age-matched WT littermates (Fig. 1F, LPS stimulation, 4- and 24-h time points), a highly significant increase in Trem1l (Fig. 1I, steady state and LPS), and a significant decrease in Trem2l (Fig. 1L, steady state and LPS).

Trem2R47H macrophages and microglia reveal a cell survival defect with defects noted primarily in apoptosis

Next, we compared the phenotypes of Trem2R47H, Trem2–/–, and WT bone marrow–derived macrophages (BMDMs) and microglia. We noted reduced proliferation and survival of the gene-edited Trem2R47H and Trem2–/– BMDMs (Fig. 2, A and B) and microglia (Fig. 2, C and D) under limiting conditions of CSF-1, consistent with previously published data on knockout alone (9, 20). As well, we noted gene dosage–dependent effects on proliferation and survival of both Trem2R47H (Fig. S2A) and Trem2–/– (Fig. S2B) myeloid cells. A more detailed analysis reveals that the variant macrophages have a primarily apoptotic and negligible necrotic response (Fig. 2, E and F) compared with the knockout macrophages, which have significant increases in both apoptotic and necrotic events (Fig. 2, G and H). Also, a breakdown of the apoptotic and necrotic events by gene dosage revealed clearly that in the case of the variant, the effects are only apparent in the homozygous R47H macrophages and not in the heterozygous macrophages, again supportive of the more subtle nature of the mutation. In the case of the heterozygous knockouts, whereas the necrotic events did not achieve statistical significance (p = 0.08), they consistently demonstrated an increasing trend compared with WT.

RNA-Seq analysis provides genome-wide molecular corroboration for the more subtle effect of the variant, including effects on several genes linked to AD

To determine whether the parallels and differences between the variant and knockout macrophages can be explained at the molecular level, we performed RNA-Seq analyses comparing WT, Trem2–/–, and Trem2R47H macrophages. At day 7 under limiting conditions of CSF-1, consistent with the gradation in severity of phenotypes observed, we observed the greatest differential transcript expression in the Trem2–/– macrophages with an intermediate effect in Trem2R47H macrophages (heat map in Fig. 3A). Pathway analyses point to a role for Trem2 in cell cycle/proliferation and survival, immune response and migration, and lipid and cholesterol homeostasis (Fig. 3B). Whereas the total number of genes involved in proliferation, necrosis, and apoptosis was higher in the knockout group, the R47H group had a larger number of apoptotic genes. Overall, the total number of genes that were differentially
Loss-of-function of R47H variant rescued by antibodies

expressed was higher in the knockout group compared with the R47H variant group. We confirmed differences in several genes in each of the modules, including genes linked to the proliferation/survival module as well as some known genetic factors linked to Alzheimer’s like ApoE (Fig. 3, C and D), pro-inflammatory cytokines that are up-regulated in AD like Il-1α (Fig. 3, E and F), and complement genes C1qa (Fig. 3, G and H) and C3 (Fig. 3, I and J). A temporal component is associated with the changes observed in some of the genes with an increase in gene expression followed by a decline, as in the case of C1qa and C3
(Fig. 3, G–J). We also noted changes in a host of chemokines/chemokine receptors in both the R47H and knockout groups, including a down-regulation of genes like Cx3cr1 (Fig. 4, A and B), Ccr2 (Fig. 4C), and Ccl2 (Fig. 4D). In a few instances, we also noted up-regulation of genes, including Ccl5 and Ccl22 (Fig. S3, A–D). The effect on Cx3cr1 was the most profound, with a significant gene dosage–dependent effect at multiple time points that was observed in both genotypes. The effects on Ccr2 and Ccl2 were more subtle and noted only at specific time points. For the first time, we noted a reduction in Flt1 in the TREM2 knockouts but no obvious effect in the variants (Fig. S3, G–J). Overall, in the majority of genes, the differential effect was significantly more pronounced in the knockouts compared with the R47H cells. The modulation of multiple migratory chemokine/chemokine receptors translated to a net reduction in migration/motility of R47H and knockout macrophages (Fig. 4E and Fig. S3G) and to a lesser extent Arg-47 microglia (Fig. 4F, trend but not statistically significant).

**In vivo LPS and in vitro Aβ challenge reveal a surprising decrease in pro-inflammatory cytokines in microglia and macrophages**

We next determined the effect of an acute *in vivo* challenge on variant and knockout microglia. Animals were administered LPS (5.0 mg/kg intraperitoneally (i.p.)), and CD11b+ microglia cells were isolated following overnight treatment according to the manufacturer’s protocols. No significant differences were observed in the total number of microglial cells isolated under homeostatic and challenge conditions (data not shown). Contrary to our expectation of increased pro-inflammatory cytokines upon LPS challenge in the variant microglia based on the proposed functional role for TREM2, we observed statistically significant decreases in pro-inflammatory cytokines, including Il-1β, Il-6, and Tnf-α (Fig. 5, A–C). In *in vitro* Aβ(1–42) (Anaspec AggreSure™) challenge of macrophages revealed a similar trend, with statistically significant decreases in CCL2, CXCL10,
and CCL5 protein levels (Fig. 5, D–F) in R47H macrophages compared with WT. A decreasing trend in CCL3 and CCL4 was also noted over and above what was observed with control scrambled A/H9252(1–42) peptide (data not shown). Similar but less significant trends were observed when microglia were treated with A/H9252(1–42), with the most significant changes observed for CCL3 and CCL4 (Fig. S4, A and B). In these instances where we measured protein levels in the conditioned media, an equal number of cells were plated at the start of the experiment, and the cells were grown in complete medium, where similar proliferation curves have been observed for WT and R47H microglia. Hence, it is unlikely that the differences observed can be attributed solely to reduced cell numbers. Further, in at least a few instances, we observed opposite trends for chemokines/cytokines, depending on the challenge administered, supportive of a generally dysregulated inflammatory state conferred by the variant/knockout that manifests differently based on the challenge. Of note, Ccl2 was one of the only chemokines tested that maintained a similar trend with most challenges. Ccl2 was reduced in Trem2−/− and Trem2R47H day 5 BMDM cultures compared with WT, although only the R47H data reached statistical significance (Fig. 4D). A small reduction of CCL2 was also noted in lavage fluid of the peritoneal cavity of knockout mice treated with zymosan (Fig. S4C) and to a lower extent (∗, p < 0.05) in the brains of LPS-treated knockout mice (Fig. S4D).

Antibodies that boost TREM2 signaling also modulate myeloid cell functioning

Whereas the Trem2R47H macrophages phenocopy (albeit to a more subtle degree) the Trem2−/− macrophages, the phenotypes cannot be explained simply by a reduction in cell-surface expression of R47H TREM2. WT and Trem2R47H BMDMs appeared to have comparable levels of surface TREM2.

Figure 3. RNA-Seq analysis provides molecular corroboration for the more subtle effect of the variant and modulation of several genes linked to AD. A, transcript heat map of differentially regulated genes between various treatment groups (n = 5 animals/genotype) is supportive of an intermediate perturbation in the Trem2R47H BMDMs compared with Trem2−/− and WT BMDMs. B, WGCNA analysis resulted in the identification of five modules/gene networks that are differentially regulated in the knockout and variant macrophages compared with WT. C–I, a subset of genes, including Apoe (C and D), Il-1α (E and F), C1qa (G and H), and C3 (I and J), confirmed as differentially regulated in R47H macrophages (C, E, G, and I; blue graphs) and knockout macrophages (D, F, H, and J; red graphs) by qPCR in a time course study. Data are presented as mean ± S.D. (error bars) and are from a representative experiment. The experiment was conducted twice independently. *, p < 0.05; **, p < 0.01; ****, p < 0.0001.
expression (Fig. S5, A and C). However, we and others have shown that the R47H variant has impaired ligand sensing/signaling (8, 21).

To determine whether the defects in proximal signaling and myeloid cell function can be rescued by a pharmacologic agent, we tested commercial and internally generated mouse antibodies in pSyk activation assays and on more distal myeloid cell functioning. We first confirmed the specificity of a commercially available TREM2 antibody (henceforth denoted by antibody 1) by demonstrating a lack of FACS shift compared with isotype control on Trem2−/− BMDMs (Fig. S5B). Next, we demonstrated that the antibody increased pSyk levels in both R47H and knockout macrophages (trend for Ccl2) at different time points. The experiment was conducted three times independently. E, migration defects were observed for the R47H macrophages in a transwell assay with C5a as the chemoattractant (data from three animals). *, p < 0.05; ***, p < 0.001; ****, p < 0.0001; data represented as mean ± S.D.
We then determined that the antibody improved survival of Trem2<sup>R47H</sup> microglia and macrophages (Fig. 6, C and D, green), whereas an equivalent rescue in cell survival was not observed when treated with isotype control antibodies (Fig. 6, C and D, blue). The effect was equally robust when we treated macrophages from aged animals (Fig. 6E). In fact, the antibody was also able to boost the function of WT macrophages and microglia (Fig. S6, A and B) but had no effect on knockout macrophage survival (Fig. S6C). Additionally, the antibody increased migration of microglia toward recombinant C5a (a classic macroglial chemoattractant) both with and without aggregated Aβ(1–42) (a putative endogenous ligand) (Fig. 6, E and F). Aβ(1–42) alone did not serve as a chemoattractant in the transwell migration assay (data not shown). It should be noted that the boost in migration reached statistical significance for the R47H microglia only in the presence of Aβ (Fig. 6G, blue bars), whereas the WT microglia showed significantly increased migration under both treatment paradigms (Fig. 6, F and G, black bars). It is also noteworthy that CCL2, a myeloid chemoattractant and the one chemokine whose level was reduced in different treatment conditions, actually increased upon antibody treatment in the Trem2<sup>R47H</sup> macrophages (Fig. S7, A and B). Additionally, antibody 1 also significantly modified expression levels of genes in module 1 (i.e., the proliferation and survival module) (>50 based on their annotation in the Gene Ontology network) in macrophages. Representative data are shown for Melk, Nek2, and Mmp14 (Fig. S7, C–E). The modulated gene signature is consistent with the antibody suppressing cell death events and promoting proliferation. To determine whether the ability to modulate different aspects of myeloid cell function was unique to antibody 1, we tested another internally generated mouse TREM2 antibody (henceforth referred to as antibody 2) that did not compete with antibody 1 for TREM2 binding and also activated Syk signaling. In this instance, antibody 2 recognized only mouse TREM2. We confirmed by FACS (Fig. S5, C and D) and immunoblotting (Fig. S5E) that antibody 2 did not have any signal in the knockout BMDMs but was able to recognize TREM2 in WT and R47H BMDMs. Antibody 2 was also able to boost survival of both R47H macrophages and microglia (Fig. 7, A and B). Further, as in the case of antibody 1, antibody 2 was also able to boost survival of WT cells (Fig. S6, D and E) but did not impact the survival or rescue survival defects observed in Trem2<sup>–/–</sup> cells (Fig. S6F). Also, the antibody did not affect migration in any of the genotypes (Fig. 7, C and D). Thus, even a comparison of just two antibodies with putatively different TREM2-binding regions and different properties resulted in different myeloid cell function modulation profiles.

**Discussion and conclusions**

In summary, we have generated powerful and specific TREM2 animal models to elucidate the effect of the R47H variant on myeloid cell function and show that pharmacologic agents like antibodies can rescue the loss of function conferred by the variant and even boost WT function. We demonstrate that targeting the Trem2 gene using gene editing allows for specific perturbation of the gene without disrupting other genes in the locus. Both TREM1 and TREM2 signal via DAP12 and probably opposing effects with respect to modulating...
inflammatory response (pro-inflammatory versus anti-inflammatory) (22); similarly, TREML1 and TREML2 also modulate inflammatory response; in fact, TREML2 is expressed on similar cells as TREM2, including microglia, and recently has been proposed to play an opposing role to TREM2 (23, 24). Hence, we expect that perturbation of multiple genes in the Trem locus as in the Trem2<sup>KOMP−/−</sup> mice can confound some of the results from mice generated by crossing these knockouts with disease models. Our TREM models will probably not suffer from these complications.

R47H variant macrophages and microglia reveal survival and migration defects in culture similar to but less severe than knockout cells (8). We note that the difference in severity in the phenotype (under challenge conditions) between a complete loss-of-function of R47H variant rescued by antibodies

Loss-of-function of R47H variant rescued by antibodies

Figure 6. An antibody that boosts TREM2 signaling also modulates myeloid cell function. A and B, antibody 1 activates TREM2/DAP12-mediated Syk signaling in Trem<sup>2<sup>−/−</sup></sup> and Trem<sup>2<sup>KOMP−/−</sup></sup> BMDMs. The effect is less pronounced in the Trem<sup>2<sup>KOMP−/−</sup></sup> BMDMs compared with WT BMDMs; the TREM2 agonist antibody does not increase pSyk levels in the Trem<sup>2<sup>−/−</sup></sup> BMDMs, confirming a TREM2-specific effect. No significant differences were observed in total Syk levels or in the loading control (actin). Data are shown from a representative set of animals. C, the antibody also restores the survival defect in Trem<sup>2<sup>KOMP−/−</sup></sup> microglia, as demonstrated in a real-time cell confluence assay. Isotype control antibody (blue) is not able to achieve the same boost in survival. Data are plotted as mean ± S.D. (error bars) and are from a single representative experiment. The experiment was conducted twice independently. D, improved survival was also noted in Trem<sup>2<sup>KOMP−/−</sup></sup> BMDMs with antibody 1 treatment (green) relative to isotype control (blue). E, BMDMs obtained from aged (18-month-old) R47H animals also demonstrate a similar survival defect that can be rescued with agonist antibody treatment (green), whereas the isotype control has no effect (blue). F, antibody 1 treatment significantly increased migration toward recombinant C5a as a chemoattractant in Trem<sup>2<sup>−/−</sup></sup> microglia relative to isotype control. G, antibody 1 treatment significantly increased migration toward recombinant C5a + Abeta(1–42) as a chemoattractant in Trem<sup>2<sup>−/−</sup></sup> and Trem<sup>2<sup>KOMP−/−</sup></sup> microglia relative to isotype control. *, p < 0.05; **, p < 0.001; ***, p < 0.0001; data represented as mean ± S.D.
deletion of the gene versus the occurrence of the variant is recapitulated at the transcript level with similar but more significant changes in the Trem2<sup>H11002</sup>/H11002/H11002 macrophages compared with R47H macrophages. The less pronounced effect of the mutation with most effects being noted primarily in the homozygous variant cells/animals is also consistent with the identification of R47H TREM2 as a rare variant associated with a polygenic disease like AD. Our data reveal how the variant can exert a subtle but cumulative effect on myeloid cell functioning over the lifetime of the individual. The occurrence of the variant alone without any other genetic risk factors confers a loss of myeloid function for the Trem2 gene that is probably exacerbated in the context of AD. Additional risk factors, notably age as well as genetic and environmental risk factors, then probably serve as epidemiological challenges and predispose heterozygous carriers to an increased risk of disease. The subtle nature of this particular variant and a comparison with heterozygous and homozygous knockouts also suggest that in the context of AD, Trem2 and Trem2<sup>H11001</sup>A<sub>H9252</sub> microglia (purple) following treatment with antibody 2 relative to isotype control (blue) both in the presence and absence of rC5a as a chemoattractant yield a <i>p</i> < 0.05; ***,<i>p</i> < 0.001; ****,<i>p</i> < 0.0001; data represented as mean ± S.D.

Pathway analyses of the differentially regulated genes provide mechanistic insight into the observed phenotypic defects. We note a dysregulation in different aspects of myeloid cell functioning, including DNA replication, cell cycle regulation, proliferation, cell death, chemokine/cytokine modulation, and complement pathway. We do not observe significant changes in genes directly linked to phagocytosis. Our data support the hypothesis that a loss of function of the TREM2 gene contributes directly to a fundamental proliferation/survival deficit that can then translate indirectly to functional effects like reduced phagocytic capacity. The role of TREM2 as a more fundamental regulator of the homeostatic state of microglia is still unclear. Recent work from Keren-Shaul et al. (25), employing single-cell RNA-Seq techniques, suggests that microglia surrounding plaques are modulated in two waves, a primary TREM2-independent wave with an up-regulation of ApoE and a down-regulation of genes like Cx3Crl and P2ry12 and a second TREM2-dependent wave that modulates lipid sensing and phagocytosis. However, some studies have reported a TREM2 depletion–linked reduction in ApoE and locking of microglia in a homeostatic state (26). Based on our gene networks and gene analysis, we find that some of the key wave 1 genes are also significantly regulated by the variant (increased ApoE, decreased Cx3Crl) under some challenge conditions. Our data suggest that TREM2 can more fundamentally regulate the transition out of a homeostatic state for myeloid cells in a challenge-dependent manner. With respect to regulation of the chemokine environment, the variant clearly contributes to a dysregulation of the chemokine environment in <i>vitro</i>. However, the observed decreasing trends in traditional pro-inflammatory cytokines upon LPS/zymosan challenge in <i>vitro</i> in variant microglia is reflective of the highly complex, temporally regulated, and
ligand/challenge-sensitive nature of the TREM2-dependent response (27–30). In fact, recent data pointing to opposing roles for TREM2 in tauopathy can potentially be attributed to the age of the animals and the different genetic models of tauopathy utilized in the two studies and further support the complex nature of TREM2-mediated modulation of neuroinflammation (15, 16).

The ability of antibodies that boost proximal signaling to rescue the viability, proliferation, and migration defects elegantly demonstrates the potential application of pharmacologic agents in rescuing cumulative genetic defects and boosting microglia activity in a TREM2-dependent manner. In fact, the ability of antibodies 1 and 2 to boost WT microglia function under challenge conditions in vitro is supportive of the putative therapeutic potential of a TREM2 agonist antibody for the >99% of LOAD patients who do not carry the R47H mutation. However, the different effects on migration, depending on which antibody is used, are supportive of more divergent and complex distal biology. The next steps will entail in vivo administration of antibodies with different in vitro profiles in neurodegenerative disease mouse models to more rigorously define the properties of an efficacious TREM2 therapeutic for the treatment of AD or other neurodegenerative diseases. Finally, most efforts on TREM2 have been focused on its functioning in microglia. Recent studies on the CNS immune environment have focused on the cross-talk between activated microglia and reactive astrocytes (31). The increase in complement genes and factors like IL-1α in the TREM2 variants and knockouts that have been proposed to play a role in modulating microglia–astrocyte cross-talk also points to potentially novel non-cell-autonomous roles for TREM2 in modulating the CNS immune environment that will need further investigation.

Materials and methods

All animal procedures were approved by the Amgen Institutional Animal Care and Use Committee.

Gene-edited in-house mouse construction

A pair of mRNAs targeting exon 2 of the TREM2 gene was synthesized. The target sequences were as follows: left, 5′-TCC-TTGAGGCGTCTCATGTAC-3′; right, 5′-TGCGCTCTCCCCAGTGCTTC-3′. The binding sites were separated by a 12-bp spacer region. A 143-mer R47H single-stranded oligonucleotide (ssODN), which was silently mutated with code modification, was synthesized. The target sequences were as follows: left, 5′-TCC-TTGAGGCGTCTCATGTAC-3′; right, 5′-TGCGCTCTCCCCAGTGCTTC-3′. The binding sites were separated by a 12-bp spacer region. A 143-mer R47H single-stranded oligonucleotide (ssODN), which was silently mutated with code modification, was synthesized. The ssODNs were ligated into the pDONR221 destination vector (Invitrogen) and then cloned into the destination vector. The target sequence in the ssODN was 5′-TCC-TTGAGGCGTCTCATGTAC-3′.

Microinjection

Two TREM2 gene-edited mRNAs were injected into the pronuclei of fertilized oocytes obtained from superovulated females of the C57BL/6 strain for TREM2 knockout. Two TREM2 gene-edited mRNAs and a 143-mer ssODN with an R47H mutation were injected into the pronuclei of fertilized oocytes obtained from superovulated females of the C57BL/6 strain for TREM2 R47H mutation. Genotyping for TREM2 KO and R47H KI mice was confirmed by PCR from tail genomic DNA.

LPS administration

4–5-month-old male KOMP and gene-edited mice were all maintained in C57BL/6 background. They were housed at a constant ambient temperature of 21 °C with a 12-h/12-h light dark cycle and were given access to food and water ad libitum. Equal numbers of animals from each genotype were randomly assigned to different groups. Treatment groups were given Escherichia coli LPS (Sigma-Aldrich) prepared in sterile saline and administered i.p. by single injection at a dose of 5 mg/kg, whereas control groups received saline i.p. Animals were dosed by a trained technician who was blinded with respect to animals’ identifications, including genotypes. Animals with clinical signs of pain or distress after treatments, including hunched posture, rough hair coat, increased respiration, and lethargy were excluded by humane euthanasia. Tissue samples were collected at the 4- and 24-h time points. Mice were euthanized with CO2 inhalation for 2 min, and blood was withdrawn by cardiac puncture. Brains were removed, divided in half, and stored at −80 °C until use. The treatment group sizes were as follows: Trem2+/− saline-treated, n = 3; WT littermates, n = 6; Trem2+/− LPS-treated, n = 3; WT littermates, n = 6; Trem2R47H+/- saline-treated, n = 5; WT littermates, n = 6; Trem2R47H−/− LPS-treated, n = 6; WT littermates, n = 6; Trem2R47H+/- saline-treated, n = 4; WT littermates, n = 5. Post hoc analysis showed that a sample size of n = 5 was adequate to detect 20% difference between groups with respect to -fold change of transcripts. A second experiment was run with Trem2+/− and Trem2R47H+/- mice (n = 6 animals in each treatment group), confirming the results shown (data not shown).

Brain qPCR

Frozen brain halves were homogenized in RNA lysis buffer, and RNA was isolated by using the RNaseasy minikit (Qiagen, Redwood City, CA). Total RNA was quantified on a NanoDrop Spectrophotometer (Thermo Fisher Scientific), and the cDNA was synthesized using the Cells-to-Ct Bulk RT Reagents (Applied Biosystems). qPCR was done in a 384-well format using TaqMan® Universal PCR Master Mix (Applied Biosystems) on a Vii7 real-time PCR system (Thermo Fisher Scientific) with the following cycle conditions: 2 min at 50 °C, 10 min at 95 °C, 40 cycles of 15 s at 95 °C, 1 min at 60 °C.

Antibodies

Rat monoclonal anti-human/mouse TREM2 (rat IgG2b clone 237920, R&D Systems) and internally generated mouse monoclonal anti-mouse TREM2 were used to activate TREM2 signaling along with the respective isotype controls (monoclonal rat IgG2b clone 141945, R&D Systems; internally generated isotype control).

Differentiation of mouse BMDMs

Bone marrow cells from Trem2−/−, Trem2+/−, Trem2R47H+/-, Trem2R47H+/+, and Trem2+/-+ mice were obtained from femurs
Loss-of-function of R47H variant rescued by antibodies

and tibiae using standard protocols (32) and seeded in DMEM supplemented with 10% heat-inactivated FBS and 50 ng/ml mouse CSF-1 (R&D Systems) in non-TC-treated sterile Petri dishes (Thermo Fisher Scientific). Nonadherent cells were removed by washing with cold PBS (Gibco) on day 5 or day 6 in culture (unless otherwise indicated), and the adherent macrophages were harvested by gentle scraping, counted, and used for further downstream analysis.

BMDM survival studies and flow cytometry analysis

Equal numbers of freshly harvested bone marrow cells from Trem2<sup>−/−</sup>, Trem2<sup>+/-</sup>, Trem2<sup>R47H</sup>, Trem2<sup>R47H/+</sup>, and Trem2<sup>++</sup> mice (n = 3 animals/genotype with the exception of WT age-matched littermate controls for day 6 samples in the knockout experiment) were differentiated and harvested on the indicated days, and cell numbers were determined using a ViCell Cell Counter (BD Biosciences). Day 6 and day 7 BMDMs were harvested and stained with FITC-annexin V and PI and analyzed on an LSR II flow cytometer (BD Biosciences). TREM2 surface expression was also measured in the macrophages using APC-conjugated mouse anti-human TREM2 antibody (clone 23790, R&D Systems).

Microglia isolation and survival study

Young mice (n = 3/genotype) were euthanized via CO<sub>2</sub> asphyxiation, and brains were dissected out. Single-cell suspensions were isolated using Miltenyi Biotec’s adult mouse brain dissociation kit, followed by microglia isolation using Miltenyi Biotec’s CD11b microbeads. Microglia were plated on a poly-D-lysine–coated 96-well plate in 200 μl of complete medium (DMEM/F-12 medium supplemented with 10% heat-inactivated FBS, 5% L929 conditioned medium, 1% penicillin/streptomycin, 1× Glutamax, 20 ng/ml granulocyte/macrophage colony–stimulating factor, 20 ng/ml M-CSF, 5 ng/ml TGF-B1). On the third day after plating and every 2–3 days thereafter, medium was replaced with basal medium (DMEM/F-12 with 10% heat-inactivated FBS). Confluence was measured using Incucyte Zoom (Essen Bioscience). For antibody treatment, basal medium was supplemented with 300 nM TREM2-activating antibodies or respective isotype control.

RNA-Seq data analysis

RNA-Seq sequencing reads were aligned using OSA aligner (35) embedded in the Omicsoft ArrayStudio pipeline (Omicsoft Inc.). Mouse genome version GRCm38 and UCSC gene annotation were used in the alignment and quantification. Quantification was performed to the gene and transcript level based on RSEM (36). Normalized gene expression level was calculated by fragments per kilobase per million reads (FPKM) and then quantile-normalized at 70 percentile to 10 (FPKQ). Only genes with at least one sample expressed at FPKQ ≥ 1 were used in the following statistical analysis. Raw read counts from the selected genes were compared using the R Bioconductor package DESeq2 following negative binomial distribution (37). Genes with Benjamini–Hochberg–corrected p value <0.05 and -fold change ≥1.5 or ≤0.5 were selected as significantly differentially expressed genes. Pathway analysis was performed using Ingenuity Pathway Analysis (IPA; Qiagen). Differentially expressed genes in both Trem2<sup>−/−</sup> versus WT and Trem2<sup>R47H</sup> were used to construct a gene co-expression network. Weighted gene co-expression network analysis (WGCNA) was performed using the Bioconductor package WGCNA. Modules in the network were defined by dynamic tree cutting in WGCNA. Genes with an absolute correlation coefficient of >0.95 with other genes were shown on the network. The red nodes are genes up-regulated in both Trem2<sup>−/−</sup> and Trem2<sup>R47H</sup>, whereas the green nodes are genes down-regulated in both Trem2<sup>−/−</sup> versus WT and Trem2<sup>R47H</sup>.

RNA-Seq confirmation of subset of differentially regulated genes

BMDMs from Trem2<sup>R47H</sup>, Trem2<sup>−/−</sup> and respective WT littermates were harvested daily between day 4 and day 8. Microglia were isolated from Trem2<sup>R47H</sup> and WT littermates treated with saline or LPS, as described previously. Total RNA was isolated using the RNeasy minikit (Qiagen) according to the manufacturer’s protocol. cDNA was generated using the RT reagents (Thermo Fisher Scientific). Primers specific to Apoe, Il-1α, Cx3cr1, C1qα, Ccl5, Ccl22, Ccr2, Flt-1, and C3 (macrophages) and Il-1β, Il-6, and Tnf-a (microglia) were ordered from Thermo Fisher Scientific, and quantitative RT-PCR was per-
Loss-of-function of R47H variant rescued by antibodies

formed in a Viia7 real-time PCR machine (Thermo Fisher Scientific) using the Taqman gene expression master mix.

For antibody treatment of macrophages before qPCR, day 5 BMDMs from Trem2\(^{+/+}\) and Trem2\(^{R47H}\) and respective WT littermates (\(n = 3\) animals/genotype) were harvested and reseeded in 6-well plates at 2 million cells/well. The cells were treated with rat IgG2b or anti-Trem2 antibody (R&D MAB17291) overnight. The subsequent steps for performing qPCR were the same as described above, using primers specific to Melk, Nek2, Mmp14, and Ccl2.

Macrophage plug area migration assay

Day 5 BMDMs from Trem2\(^{+/+}\), Trem2\(^{R47H}\), and Trem2\(^{−/−}\) mice were harvested and seeded into Radius\(^{TM}\) 96-well migration assay plates (Cell Biologics) in complete RPMI medium supplemented with 50 ng/ml M-CSF (R&D Systems). The cells were treated with either anti-TREM2 antibody, isotype control, or vehicle for 24 h. The cells were washed the next day following the manufacturer’s protocol to remove the Bio-compatible Gel layer and expose the cell-free area for migration. The medium was replaced with fresh growth medium supplemented with 50 ng/ml M-CSF and antibody, isotype, or vehicle control as above. The cell confluence was monitored using the Incucyte Zoom imaging system, and data were plotted as percent confluence.

Peritoneal lavage

Zymosan A (Sigma-Aldrich Z-4250, Saccharomyces cerevisiae) powder was suspended in PBS (1 mg/ml), sonicated with heat, and vortexed every 10 min for 30 min. The Zymosan suspension was further diluted to 0.2 mg/ml in PBS, and 500 \(\mu l\) was injected i.p. into Trem2\(^{+/+}\) and Trem2\(^{−/−}\) mice (\(n = 6\)/genotype). 3.5 and 6.5 h postinjection, mice were euthanized via CO2 asphyxiation. Peritoneal lavage was performed with 5 ml of ice-cold PBS. The peritoneal lavage fluid was centrifuged to pellet cells, and the supernatant was collected for further analysis.

Chemokine Luminex\(^{TM}\) assay

Microglia was isolated as described previously and plated on a poly-d-lysine−coated 96-well plate in 200 \(\mu l\) of complete medium at 40,000 cells/well. After 4 days, medium was replaced with complete medium supplemented with TREM2 antibody or corresponding isotype control. After 6 h, cells were treated with AggreSure\(^{TM}\) A\(\beta\)(1–42) or scrambled control (Anaspec). Final concentrations of antibody and A\(\beta\) were 300 nm and 10 \(\mu g/ml\), respectively, in a 200-\(\mu l\) volume. 16 h after A\(\beta\) treatment, cell culture supernatant was collected and assayed for chemokine levels with Chemokine 9-Plex Mouse ProcartaPlex\(^{TM}\) Panel 1 (Thermo Fisher) and analyzed with a Bio-Plex\(^{TM}\) 200 analyzer (Bio-Rad).

Chemokine ELISAs

Day 5 BMDMs from Trem2\(^{R47H}\) mice and WT littermates (\(n = 3\) animals/genotype) were harvested, and an equal number of cells were plated in 96-well plates and allowed to adhere overnight. On the following day, cells were treated with 10 \(\mu g/ml\) AggreSure\(^{TM}\) A\(\beta\)(1–42) (Anaspec) for 2 h. The conditioned medium was collected after 2 h, and chemokine levels were measured using Quantikine ELISA kits (R&D Systems). CCL2 levels in brain homogenates from LPS-treated animals (see “LPS administration” above) and peritoneal lavage from Zymosan-treated animals were also measured using the same kit. Samples were measured in triplicate. For measurement of CCL2 protein following antibody treatment, day 6 BMDMs from Trem2\(^{R47H}\) and WT littermates (\(n = 2\) animals/genotype) were treated with anti-TREM2 antibody or isotype control for 24 h in triplicate, and the CCL2 levels in the conditioned medium were measured using the same kit.

BMDM imaging

Day 6 BMDMs from Trem2\(^{+/−}\) and Trem2\(^{R47H}\) mice (\(n = 2\) animals/genotype) were harvested, and equal numbers of cells were seeded in 96-well plates. 2 h post-plating, the cells were treated with R&D monoclonal antibody or isotype control for 14 days, and cell viability was measured using CellTiter Glo (Promega). Each treatment was done in triplicate.

Cell viability

Day 6 BMDMs from 18-month-old Trem2\(^{R47H}\) and age-matched WT littermates (R47H, \(n = 22\) animals; WT, \(n = 24\) animals) were harvested, and an equal number of cells were seeded in 96-well plates. 2 h post-plating, the cells were treated with R&D monoclonal antibody or isotype control for 14 days, and cell viability was measured using CellTiter Glo (Promega).

Syk phosphorylation assay

For measurement of Syk phosphorylation, day 5 BMDMs from Trem2\(^{+/−}\), Trem2\(^{R47H}\), and respective WT littermates (\(n = 3\) animals/genotype) were plated in 6-well tissue culture−treated dishes and allowed to adhere overnight. Cells were stimulated with anti-TREM2 antibodies or appropriate controls and incubated for 10 min at 37 \(^{°}\)C. Each treatment was done in duplicate. At the end of the incubation time, the medium was removed, and cells were lysed with M-PER lysis buffer (Thermo Fisher Scientific) supplemented with HALT protease and phosphatase inhibitor (Thermo Fisher Scientific).

pSyk Western blotting

15 or 30 \(\mu g\) of total protein from the aforementioned BMDM cell lysates were separated by 4−12% BisTris SDS-PAGE (Thermo Fisher Scientific) and then transferred to polyvinylidene difluoride membranes. The membranes were probed with rabbit anti-pSYK (Cell Signaling) or rabbit anti-SYK (Cell Signaling) and mouse anti-actin (Sigma-Aldrich). Bands were visualized and quantified on the Bio-Rad imager.

Adult microglia transwell migration assay

Microglia were plated in complete medium at 5000 cells/well in transwell plates (Essen Bioscience) precoated with 20 \(\mu g/ml\) Protein G (Life Technologies) and 5 \(\mu g/ml\) I-CAM (Life Technologies). Transwells were placed in reservoir plates containing 200 \(\mu l\) of complete medium with 1 \(\mu g/ml\) recombinant C5a (R&D Systems). For migration assays with antibody treatment, microglia were plated at 3000 cells/well in complete medium.
supplemented with 300 nM TREM2 antibodies or corresponding isotype control. Transwells were initially placed in a reservoir with complete medium, and on day 5, medium in the transwell was replaced with 300 nM TREM2 antibodies or corresponding isotype controls in complete medium. Transwells were moved to a new reservoir plate containing 200 μl of complete medium with 1 μg/ml recombinant C5a and 100 ng/ml recombinant C5a and 10 μg/ml AggreSure™ Aβ1–42 (Anaspec). Transwell migration was monitored and analyzed using Incucyte Zoom (Essen Bioscience).


**References**


