Multiple sclerosis (MS) is an inflammatory disease of the CNS that is characterized by BBB dysfunction and has a much higher incidence in females. Compared with other strains of mice, EAE in the SJL mouse strain models multiple features of MS, including an enhanced sensitivity of female mice to disease; however, the molecular mechanisms that underlie the sex- and strain-dependent differences in disease susceptibility have not been described. We identified sphingosine-1-phosphate receptor 2 (S1PR2) as a sex- and strain-specific, disease-modifying molecule that regulates BBB permeability by destabilizing adherens junctions. S1PR2 expression was increased in disease-susceptible regions of the CNS of both female SJL EAE mice and female patients with MS compared with their male counterparts. Pharmacological blockade or lack of S1PR2 signaling decreased EAE disease severity as the result of enhanced endothelial barrier function. Enhanced S1PR2 signaling in an in vitro BBB model altered adherens junction formation via activation of Rho/ROCK, CDC42, and caveolin endocytosis-dependent pathways, resulting in loss of apicobasal polarity and relocation of abluminal CXCL12 to vessel lumina. Furthermore, S1PR2-dependent BBB disruption and CXCL12 relocation were observed in vivo. These results identify a link between S1PR2 signaling and BBB polarity and implicate S1PR2 in sex-specific patterns of disease during CNS autoimmunity.
Enhanced sphingosine-1-phosphate receptor 2 expression underlies female CNS autoimmunity susceptibility

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Multiple sclerosis (MS) is an inflammatory disease of the CNS that is characterized by BBB dysfunction and has a much higher incidence in females. Compared with other strains of mice, EAE in the SJL mouse strain models multiple features of MS, including an enhanced sensitivity of female mice to disease; however, the molecular mechanisms that underlie the sex- and strain-dependent differences in disease susceptibility have not been described. We identified sphingosine-1-phosphate receptor 2 (S1PR2) as a sex- and strain-specific, disease-modifying molecule that regulates BBB permeability by destabilizing adherens junctions. S1PR2 expression was increased in disease-susceptible regions of the CNS of both female SJL EAE mice and female patients with MS compared with their male counterparts. Pharmacological blockade or lack of S1PR2 signaling decreased EAE disease severity as the result of enhanced endothelial barrier function. Enhanced S1PR2 signaling in an in vitro BBB model altered adherens junction formation via activation of Rho/ROCK, CDC42, and caveolin endocytosis-dependent pathways, resulting in loss of apicobasal polarity and relocation of abluminal CXCL12 to vessel lumina. Furthermore, S1PR2-dependent BBB disruption and CXCL12 relocation were observed in vivo. These results identify a link between S1PR2 signaling and BBB polarity and implicate S1PR2 in sex-specific patterns of disease during CNS autoimmunity.

Introduction

Multiple sclerosis (MS) is an autoimmune, demyelinating disease of the CNS that has a strong sex bias, with the female to male ratio currently ranging from 3:1 to 4:1 (1–3). Relapsing-remitting MS (RRMS), the most common form of the disease in women and men, is a condition in which recurrent relapses of new neurological dysfunction (relapses) are separated by periods of clinical stability. The mechanisms underlying sex differences in MS and whether they predominantly affect immune responses, CNS susceptibility to inflammation, or both are unclear.

EAE in the inbred SJL mouse strain is commonly used to model the sexual dimorphism observed in MS. Female SJL mice exhibit increased sensitivity to EAE and a relapsing-remitting phenotype, while male SJL mice exhibit monophasic disease resembling EAE induced in both sexes of other mouse strains, such as C57BL/6. Gonadal hormones and epigenetic regulation of sex chromosomes are postulated to contribute to sex differences in SJL mice during EAE and other autoimmune disorders (4, 5). Several studies also indicate that the CNS itself may be responsible for the observed sex differences in disease expression, primarily via alterations in endothelial cell-mediated regulation of immune cell entry (6–9).

Studies of endothelium in peripheral organs implicate cadherins and sphingosine-1-phosphate (S1P) signaling in mediating alterations in cadherin-dependent barrier properties between endothelial cells (10). Cadherins are plasma membrane proteins associated with adherens junctions (AJs), whose expression at cell-cell contacts depends on endocytic transport (11–13). Vascular endothelial–cadherin (VE-cadherin) expression is required for CNS endothelial polarity and vascular lumen organization (14), and cytokine-mediated alterations in VE-cadherin expression in CNS endothelium influence leukocyte entry (15, 16). S1P, a bioactive metabolite of sphingolipids, is produced by erythrocytes in the plasma, vascular and lymphatic endothelial cells, and neuronal lineage cells within the CNS (17–19). S1P signals via 5 G protein-coupled receptors (S1PRs) to regulate various physiological responses, including vascular permeability (20–22). Endothelia express different combinations of S1P receptors (S1PRs), which regulate endothelial cell survival, migration, AJ assembly, and barrier integrity (20, 21, 23). S1PRs have received considerable attention in the MS field due to recent success with the broad S1PR inhibitor, fingolimod (FTY720-phosphate), which binds to S1PR subtypes S1PR1, S1PR3, S1PR4, and S1PR5 but not S1PR2 (24), and inhibits vascular endothelial cell growth factor–induced vascular permeability in vivo (22). Several in vitro studies demonstrate roles for S1PR2 in vascular biology, including regulation of portal vein pressure (25), the formation of atherosclerotic plaques (26), inflammation (27), and retinal angiogenesis (28). In vitro studies implicate S1PR2 in the regulation of vascular integrity via phosphorylation of VE-cadherin, preventing its translocation to cell-cell contact sites (29). The expression or function of S1PR2 at endothelial barriers within the CNS or during induction of CNS autoimmunity has not been investigated.

Here, we identify a sexually dimorphic target for the treatment of relapsing-remitting CNS autoimmunity. Microarray analysis of specimens from naive male and female adult SJL mice revealed...
The inbred SJL mouse strain has been used as a model of the sexual dimorphism observed in MS, as female SJL mice are more susceptible to EAE than male SJL mice and exhibit a relapsing-remitting disease pattern similar to that observed in patients with MS (33). We hypothesized that sexual dimorphism in this MS model arises from sexually dimorphic and CNS-region specific expression of genes that regulate BBB permeability and leukocyte entry. Thus, we used a whole transcriptome microarray to identify sexually dimorphic RNA transcripts that are differentially expressed in disease-susceptible versus nonsusceptible CNS regions of male and female SJL mice. Profiling total RNA from the cerebella and frontal cortices of naïve female SJL mice versus male SJL mice revealed differential expression of 44 and 23 gene transcripts, respectively, with 20 being unique to the cerebellum (Table 1 and Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI73408DS1). Sixteen of these transcripts have uncharacterized genetic function and are only identified by their locus link or RIKEN database numbers (Supplemental Table 1). Fifteen of these uncharacterized loci are linked to the Y chromosome, and one is autosomal. The 3 loci that were dimorphic in both cortex and cerebellum are linked to the X or Y chromosomes, including Xist, which regulates X-chromosome inactivation (Table 1). Of the 4 novel autosomal loci that were unique to the cerebellum, 2 were of interest for autoimmune disease: the IL-20 receptor b, a receptor associated with skin autoimmune diseases (34–37), and sphingosine-1-phosphate receptor 2 (S1PR2), which regulates vascular permeability in peripheral organs (refs. 29, 38, 39, and Figure 1A). Because S1PR2 belongs to a family of G protein-coupled receptors expressed by cells of the vascular, immune, and nervous systems and S1PR2 signaling, in particular, disrupts endothelial AJs, we focused our studies in CNS autoimmunity on this protein (29). Quantitative PCR (QPCR) evaluation of sexually dimorphic expression of S1PR2 in cerebella of SJL mice validated microarray results (P = 0.0255) (Figure 1B). Western blot analysis confirmed microarray and QPCR data, revealing significantly enhanced expression of S1PR2 in the cerebella of naïve female SJL mice compared with that in male SJL mice (P < 0.05), while no sexual dimorphism was observed in naive C57BL/6 mice (Figure 1C). Whole brain extracts from S1PR2-deficient mice confirmed antibody specificity (Supplemental Figure 1A). Of interest, Western blot examination of S1PR2 levels within the cerebella of naïve SJL-Chr YB10.S male mice compared with those of SJL male and female mice did not suggest that a Y-chromosome polymorphism (40) is responsible for sexually dimorphic S1PR2 expression in the CNS of SJL mice (Supplemental Figure 2).

Because S1PR2 has known roles in vascular biology, we assessed disruption of the BBB, via administration of sodium fluorescein to naïve male and female animals of both B6 and SJL backgrounds. Consistent with increased S1PR2 expression in the cerebella of naïve female SJL animals, fluorescein permeability was significantly increased in the cerebella of naïve female SJL mice compared with that in male mice (P < 0.01) (Figure 1D). Similar to wild-type B6 mice, S1PR2-deficient mice and their wild-type controls, which are both on a 129Sv/B6 background, do not exhibit sexual dimorphism and deficiency of S1PR2 does not alter BBB permeability in naïve animals (Supplemental Figure 1B). Immunohistochemical detection of extravasated albumen in CNS white matter of naïve SJL mice also exhibited sexual dimorphism (Figure 1E). Of note, sexual dimorphism in S1PR2 expression and vascular permeability was not observed in the spinal cord, which is a disease-susceptible CNS region in SJL mice. In addition, alterations in levels of claudin 5, as assessed

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**Table 1**

Sexually dimorphic genes or loci in naïve SJL cerebella and frontal cortices

<table>
<thead>
<tr>
<th>Gene symbol</th>
<th>CB (female vs. male)</th>
<th>CTX (female vs. male)</th>
<th>Chromosome</th>
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<tr>
<td><strong>Known X- or Y-linked loci</strong></td>
<td></td>
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<td></td>
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<td>3.837 a</td>
<td>2.544 a</td>
<td>X</td>
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Summary of sexually dimorphic genes or loci in SJL cerebella (CB) and frontal cortices (CTX) and chromosome locations. Four novel autosomal loci were differentially expressed in the female cerebella (shown in bold).
improved by Western blotting, did not exhibit sexual dimorphism in naive B6 or SJL mice (Supplemental Figure 3). Immunohistochemistry (IHC) analysis of cellular sources of S1PR2 in the CNS of naive female and male SJL mice revealed localization mostly along endothelial cells, with some astrocytes and pericytes as additional sources (Figure 2).

Female SJL mice and humans exhibit increased vascular S1PR2 during CNS autoimmunity. To assess sexually dimorphic alterations in BBB permeability during CNS autoimmunity, we administered sodium fluorescein to male and female animals of both B6 and SJL backgrounds after induction of EAE via immunization with appropriate myelin peptides. During the course of EAE, fluorescein permeability increased in all CNS regions in both sexes and strains, with significantly higher permeability in female SJL mice compared with that in male SJL mice at peak of disease and during remission in cortices (peak: $P < 0.05$; remission: $P < 0.05$) and cerebella (peak: $P < 0.05$; remission: $P < 0.001$) (Figure 3A). Expression of S1PR2 was also sexually dimorphic during EAE, as S1PR2 expression was increased in disease-susceptible regions (cerebella: $P < 0.05$; spinal cord: $P < 0.001$) in female SJL mice at peak of EAE, with no detectable differences in cortices (Figure 3B). Again, no such sexual dimorphism was observed in C57BL/6 mice. Although ovariectomy and estrogen exposure affect disease in female SJL mice (33, 41–45), levels of S1PR2 expression in all CNS regions of naive female SJL mice or of female SJL mice immunized with proteolipid protein epitope 139–151 (PLP 139–151) were unaffected by ovariectomy, with or without 17β-estradiol replacement (Supplemental Figure 4). Of note, sham-ovariectomized and ovariectomized mice were evaluated when they attained clinical scores of 2, whereas ovariectomized mice treated with 17β-estradiol did not develop symptomatic EAE. Thus, suppression of clinical symptoms by 17β-estradiol in PLP139-151-immunized mice did not affect CNS levels of S1PR2 expression. These data are also consistent with studies demonstrating that 17β-estradiol does not affect systemic inflammation in the setting of active immunization (41).
S1PR2 immunoreactivity was localized to endothelium, vessel-associated astrocytes, and occasional pericytes within the CNS of SJL mice with EAE (Figure 3C) and within human post-mortem cerebellar specimens derived from patients with and without MS (Supplemental Table 2 and Figure 4, A and B). Furthermore, quantitative IHC assessment of human specimens showed significantly higher levels of vessel-associated S1PR2 in patients with MS compared with that in patients without MS (P = 0.01) and a trend toward increased S1PR2 expression in female patients with MS compared with that in male patients with MS (Figure 4C). Of interest, the highest levels of S1PR2 expression in female patients with MS were in those with RRMS (2 cases), and the 1 male patient with high levels of S1PR2 had a history of occipital head trauma. These results suggest that S1PR2 may play a role in sexually dimorphic patterns of BBB dysfunction in the setting of CNS autoimmunity.

Clinical disease severity and BBB permeability during EAE are reduced by pharmacologic or genetic inactivation of S1PR2. To investigate whether S1PR2 modulates the pathogenesis of EAE, we evaluated clinical disease scores in the setting of both pharmacologic blockade and genetic deletion of S1PR2. Female SJL mice treated with the specific S1PR2 antagonist JTE-013 (1.5 mg/kg) (46), beginning when mice reached a clinical score of 2 and continuing for 10 (Figure 5A, top) or 30 (Figure 5A, bottom) days, exhibited less severe daily clinical scores (P < 0.0001, both lengths of treatment), maximum severity scores (P < 0.0001, both lengths of treatment), and mean cumulative scores (P < 0.01 for 10 days treatment, P < 0.001 for 30 days treatment), while male SJL mice did not respond to treatment (Supplemental Figure 5). No changes in body weight were observed after JTE-013 treatment, regardless of treatment duration (Supplemental Figure 6, A–C). Consistent with clinical assessments, H&E staining revealed significantly fewer inflammatory foci within the white matter of both spinal cords (P < 0.01) and cerebella (P < 0.05) of JTE-013 mice compared with that in vehicle-treated female SJL mice at peak of EAE (Figure 5B), and cerebellar lesions were significantly reduced (P < 0.05) during remission. Luxol fast blue (LFB) staining further revealed that JTE-013 treatment reduced demyelination of spinal cords and cerebellar white matter during peak and remission phases of EAE (Figure SB), and female SJL mice with EAE treated with JTE-013 for 5 days also exhibited significantly reduced fluorescein permeability in both cerebella (P < 0.001) and spinal cords (P < 0.01), compared with
vehicle controls (Figure 5C, top) and a significant reduction in spinal cord permeability after prolonged treatment for 30 days (P < 0.01) (Figure 5C, bottom). Similar results were observed via immunohistochemical detection of extravasated albumen within the white matter of diseases mice (Supplemental Figure 7A).

Consistent with results after antagonist treatment, S1PR2-deficient mice immunized with MOG<sub>35-55</sub> exhibited significantly decreased severity of EAE compared with that of wild-type controls, indicated by lower daily clinical scores (P < 0.0001), maximum severity scores (P < 0.0001), and mean cumulative scores.
**Figure 4**

BBB expression of S1PR2 is increased in MS. (A) Endothelial cell (CD31, green) and (B) astrocyte (GFAP, green) localization of S1PR2 (red) in female and male cerebellar tissue obtained from patients with and without MS. Nuclei are counterstained with Topro3 (blue). (A) Control stains, in which sections were first blocked with immunogen (1 mg/ml) prior to detection of CD31 (green) and S1PR2 (red), are included. Scale bar: 25 μm. (C) Quantification of amounts of vessel-associated S1PR2 fluorescence in female (red circles) and male (blue circles) samples from patients with and without MS. Levels of S1PR2 fluorescence were determined by examining S1PR2 staining in venule ROIs in 10 images per patient (4–6 patients per group), normalized by area of CD31 staining to control for size and numbers of vessels. Note that the outlier in the male MS group also had a history of occipital head trauma. *P < 0.05. Horizontal bars represent geometric means.

(P < 0.0001) (Figure 6A). *S1pr2*−/− animals also exhibited less demyelination and fewer inflammatory foci during peak of EAE, compared with wild-type mice, in the spinal cords (P = 0.03) but not the cerebella (Figure 6B), in addition to significant reductions in fluorescein permeability in the spinal cords during both acute (P < 0.01) and chronic (P < 0.01) EAE (Figure 6C). There was also less detection of extravasated albumen in the spinal cords of *S1pr2*−/− mice compared with that in wild-type animals at peak of disease (Supplemental Figure 7B).

S1PR2 signaling enhances BBB permeability via caveolin-endocytosis, Rho/ROCK, and CDC42-dependent pathways. To further assess the impact of S1PR2 signaling on BBB function, we used a well-characterized transwell in vitro model of the BBB consisting of human cerebral microvascular endothelial cells (HCMECs) grown on porous filter membranes over primary human astrocytes (47). Barrier integrity was evaluated by electrode recording of transendothelial electrical resistance (TEER). Treatment with exogenous S1P permeabilized in vitro BBBs in a dose-dependent manner over the course of 4 hours, indicated by significant reductions in TEER (P < 0.001, all comparisons) (Figure 7A). This disruption of barrier integrity was S1PR2 dependent, as treatment with an S1PR2 antagonist (Figure 7B), but not an S1PR1 antagonist (Figure 7C), rescued TEER reduction after S1P treatment. Moreover, treatment with an S1PR1-specific agonist led to a small increase in TEER in both vehicle- and S1P-treated cultures (P < 0.05) (Supplemental Figure 8A), in contrast to the TEER decreases produced by S1P via S1PR2. Use of RNA interference to specifically knockdown S1PR1 and S1PR2 similarly demonstrated S1PR2-dependent disruption.
of barrier integrity (Supplemental Figure 9, A and B). Further characterization of the signaling components that contribute to S1PR2 dysregulation of the in vitro BBB revealed that pharmacological blockade of the caveolin-dependent endocytic pathway (Figure 7D) prior to S1P treatment prevented TEER reductions (P < 0.001), while blockade of clathrin-dependent endocytosis (Supplemental Figure 8B) or macropinocytosis (Supplemental Figure 8C) produced either small or no changes in TEER after S1P administration, respectively. TEER reductions after S1P treatment were also prevented by pretreatment with inhibitors of the Rho GTPase effectors Rho/ROCK (P < 0.001) (Figure 7E) and CDC42 (P < 0.001) (Figure 7F) but not Rac (Supplemental Figure 8D).

S1PR2 signaling leads to breakdown of AJs and loss of polarized expression of CXCL12 on basolateral/abluminal surfaces. We next examined whether S1PR2-dependent disruption of BBB function resulted in molecular changes to the structure of the BBB endothelium. Similar to results with TEER recording, S1P treatment resulted in a breakdown of AJs, indicated by loss of intensity and discrete-ness of VE-cadherin staining at intercellular borders (Figure 7G, left). Diminished AJ integrity was also associated with a loss of polarized protein expression on the apical and basolateral surfaces of endothelial cells, as assessed via z-plane confocal microscopy (imaging on same plane of microscopy z axis) for canonical markers of apical (γ-glutamyltransferase-1 [GGT]) and basolateral (CXCL12) BBB surface markers (Figure 6G, right). Again, loss of AJ integrity and polarized GGT/CXCL12 expression after S1P treatment was specific to S1PR2, as it could be reversed by JTE-013 treatment (Figure 7H) but not by treatment with a specific antagonist (Figure 7I) or agonist of S1PR1 (Supplemental Figure 8E). Knockdown of S1PR2, but not S1PR1, RNA in the context of S1P treatment also led to loss of AJ integrity (Supplemental Figure 9C).

Similarly, pretreatment with inhibitors of caveolin-dependent endocytosis (Figure 7J) and Rho/ROCK or CDC42 (Figure 7, K and L) prevented loss of AJs and endothelial polarity, while inhibition of clathrin-dependent endocytosis, macropinocytosis, and Rac (Supplemental Figure 8, F–H) had no effect.

Given the mouse strain- and sex-specific differences in CNS levels of expression of S1PR2, we next analyzed polarized CXCL12 expression at the BBB in naïve female and male mice of various strains susceptible to EAE via quantification of fluorescence intensity across vessels using confocal microscopy (32). In the spinal cord white matter, CXCL12 expression was predominantly abluminal on the vessels of male SJL mice and in C57BL/6 and B10.PL mice of either sex, whereas vessels in female SJL mice instead exhibited a unique pattern of primarily lumenal CXCL12 expression (Supplemental Figure 10A). Similar analyses of brain vasculature in female and male SJL mice revealed a predominantly lumenal display of CXCL12 in the brainstem and cerebellar white matter tracts in the female mice only, while vessels in all brain regions of male mice and in supratentorial brain regions of female mice exhibited proper abluminal patterns of CXCL12 (refs. 31, 32, and Supplemental Figure 10B). After induction of EAE, inflamed vessels within infratentorial regions of male mice exhibited a shift in CXCL12 expression from abluminal to luminal locations, while female mice exhibited loss of CXCL12 expression (Supplemental Figure 11), as previously reported (30). To test whether inactivation of S1PR2 altered BBB expression of CXCL12 in vivo, we analyzed the location of CXCL12 at CNS vasculature within the cortices, cerebella, and spinal cords of female SJL mice treated with vehicle or JTE-013 as well as wild-type mice and S1pr2−/− mice at peak of disease. JTE-013–treated
female SJL mice exhibited significantly increased levels of abluminal CXCL12 in infratentorial CNS regions (cerebella: $P < 0.01$; spinal cords: $P < 0.05$), with no changes observed in cortical vessels (Figure 8, A and B). Similarly, $S1pr2^{-/-}$ mice exhibited significantly higher levels of abluminal CXCL12 within spinal cord white matter regions than wild-type controls at peak of EAE ($P < 0.001$) (Figure 8, C and D). These data suggest that S1PR2 activity dysregulates proper abluminal expression of CXCL12 and that this affects disease course during CNS autoimmunity.

**Discussion**

We have defined a sexually dimorphic role for S1PR2 in the disruption of polarity at the BBB. Elevated levels of S1PR2 were detected in EAE-susceptible CNS regions in female SJL mice compared with that in male SJL mice and compared with C57BL/6 mice of both sexes. Consistent with this, antagonism of S1PR2 ameliorates EAE and alterations in BBB permeability. Wild-type and $S1pr2^{-/-}$ mice were immunized with MOG and followed for disease progression. (A) Clinical EAE scores ± SEM. The inset shows mean of cumulative score and highest severity score ± SEM for 10 to 15 mice per genotype. (B) H&E-stained (left panels) and LFB-stained (right panels) sections derived from spinal cords and cerebella of WT and $S1pr2^{-/-}$ mice at peak of EAE. Scale bar: 25 μm. Bar graphs depict mean numbers of white matter lesions within both CNS regions ± SEM for 4 to 6 mice per genotype. (C) Relative BBB permeability of cortex, cerebellum, and spinal cord tissues derived from WT and $S1pr2^{-/-}$ mice at peak and chronic stages of monophasic EAE. Data are depicted as mean fluorescence intensity, normalized against plasma values for individual mice ± SEM for 4 to 6 mice per genotype, with means normalized against mean values for WT naive controls. *$P < 0.05$; **$P < 0.01$; ***$P < 0.001$.

S1PR2 deficiency ameliorates EAE and alterations in BBB permeability. Thus, analysis of S1PR2 expression in the cerebella of SJL/J-Chr YB10.5 male mice did not link sexually dimorphic effects to the Y chromosome (40), and analysis of S1PR2 expression in the CNS of naive or diseased, ovariectomized mice, with or without estradiol replacement, also did not reveal acute regulation of S1PR2 by gonadal hormones. Most current therapies for RRMS are immunomodulators, while others address the consequences of relapses by managing symptoms or improving function (51, 52). However, these drugs often have off-target side effects, because they do not specifically target myelin-reactive leukocytes (53, 54). Targeting S1PR2 may avoid common off-target effects and maybe more CNS specific than the use of FTY720, which targets several S1PRs but not S1PR2 (24). As S1PR2 also maintains germinal center B cells (55), its blockade is unlikely to lead to the lymphopenia observed with FTY720. FTY720 is also associated with severe adverse events, including fatal herpesvirus infections, hemorrhagic focal encephalitis, and vasospasm of the brachial artery, mostly mediated by S1PR3 signalling after prolonged use (56–62).

S1PR2 activity in our study was associated with several critical markers of endothelial barrier dysfunction, including higher transendothelial permeability and loss of apicobasal polarity, in vitro and in vivo. Our findings corroborate in vitro prior reports, suggesting counterbalancing roles of S1PR1 and S1PR2 in maintaining vascular endothelial integrity via Rho GTPase signaling, with S1PR1 enhancing barrier function primarily via Rac activation (20, 63, 64) and S1PR2 disrupting barrier function through cytoskeletal contraction, stress fiber formation, and junction protein disassembly via RhoA/ROCK signaling (29, 65). Our results also implicate potentially complementary roles for RhoA and CDC42, a key regulator of cellular polarity (66), in S1PR2-mediated BBB dysregulation, perhaps by convergent downstream effector signaling (67). The cavelolin-dependent endocytic pathway, but not the clathrin-dependent endocytic or macropinocytic pathways, also appears to be an important mediator of barrier dysregulation via S1PR2, likely due to the enrichment of S1PRs in close proximity with their associated effectors in membrane caveolae (64), though evidence...
S1PR2 signaling dysregulates CNS endothelial barrier structure and function through Rho/ROCK, CDC42, and caveolin endocytosis-dependent pathways. (A–F) Paracellular permeability of in vitro BBB cultures was assessed by electrode recording of TEER, reported in Ω/cm². (A) TEER after treatment of in vitro BBB cultures with 10, 100, or 1,000 nM exogenous S1P for 4 hours or (B and C) 1,000 nM S1P for 4 hours with or without (B) 1,000 nM JTE-013 at 2 hours or (C) the S1PR1-specific antagonist W146 (1,000 nM) at 2 hours. (D–F) TEER after 1,000 nM S1P treatment of BBB cultures pretreated for 2 hours with (D) the caveolin endocytosis inhibitor MBCD (10 mM), (E) Rho-associated protein kinase (ROCK) inhibitor H1152P (10 nM), or (F) CDC42 inhibitor ML141 (100 nM). TEER values are mean ± SEM of 6 to 9 replicates of 2 to 3 independent experiments. ***P < 0.001, repeated-measures 2-way ANOVA. (G–L) Immunocytochemical staining of AJs in HCMED/D3 cells via labeling of VE-cadherin (red, left and middle panels; scale bar: 30 μm) and confocal z-stack reconstruction of HCMED/D3 cells (right panels; scale bar: 15 μm), demonstrating polarized expression of canonical apical marker GGT-1 (green) and basolateral CXCL12 (red) after treatment with (G) vehicle or 1,000 nM S1P for 4 hours followed by treatment with (H) JTE, (I) W146, (J) MBCD, (K) H1152P, or (L) ML141 treatment at 2 hours. Inhibitor concentrations in H–L are identical to those in B–F. Immunocytochemical images are representative images of 2 to 3 independent experiments. IC, isotype controls.
also suggests that AJs can be disrupted via caveolin-mediated endothelial dysfunction of junction proteins (13, 68, 69). Of interest, interferon-beta, which prevents inflammatory events in ERMVM, promotes stability of VE-cadherin (70), suggesting that the specific targeting of this pathway could greatly improve therapy. Finally, S1PR2 antagonism does not affect receptor expression by endothelial cells (71), targeting the receptor to prevent alterations in BBB permeability should not lead to long-term effects on BBB function.

Association of S1PR2 with endothelial permeability by VE-cadherin–mediated tight junction dysfunction was first reported by Sanchez et al. using an in vitro model of HUVECs genetically modified to express S1PR2 (72). However, the observed amelioration of EAE in S1PR2-deficient mice could be attributed potentially to other immune mechanisms that do not incorporate effects of endothelial barriers. Among the possibilities are the effects of receptor deficiency on astrocytes or pericytes (73–76); the absence of antigen capture, as observed in vitro for Langerhans cells (77) and alveolar macrophages (78); or loss of B cell retention at germinai centers (79, 80), which could alter their ability to function as effector or antigen-presenting cells within the CNS. Last, S1PR2 signaling is required for degranulation and cytokine release by activated mast cells (81, 82), which have been implicated in the neuropathogenesis of MS and EAE (83).

We reported previously that postcapillary venules in the center of MS lesions display the normally abluminal chemokine, CXCL12, aberrantly along luminal surfaces (30–32, 84). In our study, altered patterns of CXCL12 expression at the BBB were found to be associated specifically with MS compared with other neuroinflammatory conditions, including viral encephalitis and CNS lymphoma (31). Redistributed CXCL12 was associated with increased activation of CXCR4 on leukocytes within vessel lumen, suggesting that altered CNS vasculature polarity increases leukocyte capture at endothelial cell surfaces. The finding that humans with susceptibility for MS exhibit high levels or sexually dimorphic expression of S1PR2 suggests that the molecule may underlie susceptibility to CNS autoimmunity and might be a disease-modifying target in MS, similar to that observed in the female SJL mouse. The identification of S1PR2 as a target for BBB stabilization that preserves CXCL12 polarity and therefore immune privilege is an exciting advance toward the development of novel MS therapies that limit CNS inflammation without compromising immune function.

Figure 8
In vivo S1PR2 inactivation preserves BBB polarity during EAE. Detection of CD31 (green) and CXCL12 (red) within cortices, cerebellum, and spinal cords of (A and B) female SJL mice treated with vehicle or JTE-013 (1.5 mg/kg) or (C and D) wild-type and S1pr2−/− mice, all at peak of EAE. (A and C) Nuclei have been stained with Topro3 (blue). Scale bar: 10 μm. Data are representative of approximately 30 images each from 3 to 5 mice per treatment group. Quantification of fluorescence intensity during confocal microscopy for CXCL12 (red lines) and CD31 (green lines) is shown below microphotographs. Arrows indicate location transected in line plot depictions. (B and D) Quantitative analysis of location of CXCL12 expression on CD31− venules within CNS tissues of (B) vehicle- and JTE-013–treated and (D) wild-type and S1pr2−/− mice at peak of EAE. Data are derived from venules analyzed within 30 images per brain region for 3 to 5 mice per treatment group and are expressed as (mean ± SEM) percentages of vessels with abluminal, lumenal, or absent CXCL12 signal. *P < 0.05; **P < 0.01; ***P < 0.001 for χ² comparisons between CXCL12 locations within each brain region. Figure 8

Methods

**Animals.** SJL/JOrlCrI mice (SJL, Charles River); C57BL/6, C57BL/10, and SJL/J-Chr YB10.S mice (The Jackson Laboratory); B6129SF1 and 129S6/SvEvTac mice (Taconic Farms); and S1pr2−/− mice were maintained in pathogen-free conditions (Department of Comparative Medicine, Washington University, St. Louis, Missouri, USA). The S1pr2−/− mice, which are on a B6129S background, were generated and genotyped as previously described (85). All mice were 10 weeks old when used for experiments.

**Microarray analysis.** Naive female and male SJL littmates (n = 6) were intracardially perfused with RNAlater (Ambion, Life Technologies), followed by dissection of the cerebellum and frontal cortices (2 mm). Tissue was quickly frozen in TRIzol (Ambion, Life Technologies), and RNA was extracted via standardized protocols. Transcriptional profiling was done using the Illumina Mouse4 Expression BeadChip v2 (Illumina Inc.). 100 ng total RNA was used to produce an amplified pool of biotin-labeled RNA corresponding to the polyadenylated mRNA fraction. Biotinylated RNA was hybridized to the array and stained with fluorescently labeled antibiotin antibody. Then, the BeadChip was scanned on an Illumina BeadArray Reader (Illumina Inc.). Raw data were imported into Partek Genomics Suite (Partek Incorporated) for further characterization of sexually dimorphic gene expression in female cerebella. Microarray data were deposited in GEO (accession no. GSE55718).

**Antibodies.** The following antibodies were used for IHC or Western blotting: goat anti-mouse albumin conjugated to HRP (Novus); biotinylated rabbit polyclonal anti-human CXCL12 (Peprotech); rabbit anti-S1PR2/Edg-5 and rabbit anti-S1P1/Edg-1 antibodies (Acris Antibodies Inc.); mouse anti-β-tubulin (Sigma-Aldrich); monoclonal mouse anti-human VE cadherin, monoclonal rat anti-mouse, and rat anti-human CD31 (BD Bioscience); mouse monoclonal anti-human glial fibrillary acidic protein (GFAP) and goat anti-mouse PDEGF-Rb (R&D Systems); rabbit anti-VE cadherin/phospho Y658 and rabbit anti–GFT-1 (Abcam Inc.); rabbit anti-human GFAP (Invitrogen, Life Technologies); IgG isotype antibodies (mouse and rabbit, Invitrogen, Life Technologies; rat, BD Bioscience); conjugated-secondary antibodies, streptavidin conjugates, and nuclear stains (Molecular Probes); and IRDye-conjugated secondary antibodies for Western blots (LI-COR Biosciences Inc.).

**QPCR.** Total RNA derived from CNS tissues was prepared and QPCR was performed as previously described (86). Amplification of S1PR2 was done using 5′-ATGGGGCGGCTTATACTCAGAG-3′ (sense) and 5′-GACGGA-GAACGAT GTGTCACCA-3′ (anti-sense) primers as reported by Edsbagge (87). QPCR products were validated via sequencing by the Protein and Nucleic Acid Chemistry Laboratory at Washington University.

**Western blotting for S1PR2 in murine CNS.** Protein lysates (40 μg) of CNS tissues derived from 10-week-old SJL and C57BL/6 mice of both sexes (n = 8) and after ovariec-tomy, with and without treatment with 17β-estradiol or placebo, were prepared in RIPA buffer supplemented with 1:100 protease inhibitor cocktail and 1:100 phosphatase inhibitor cocktail-3 (all from Sigma-Aldrich) and resolved with 10% Bis-Tris gels and transferred onto iBlot Nitrocellulose transfer membranes (both from Invitrogen) according to standard protocols. Blots were probed with polyclonal rabbit anti-S1PR2 antibody. Total protein loading per lane was evaluated with mouse anti-β-tubulin antibody. This was followed by incubation with IRDye-conjugated secondary antibodies (LI-COR). Blots were imaged with the Odyssey fluorescent scanning system (LI-COR).

**Ovariectomy and hormone replacement in female SJL mice.** Female SJL mice were ovariec-tomized at 6 weeks of age mice prior to sexual maturity via standard protocols (33). Briefly, anesthetized animals underwent a 1-cm transverse midlumbar skin incision, followed by blunt dissection of underlying musculature. The ovarian fat pad was exteriorized, ovarian vessels were ligated, and ovaries were excised, followed by skin closure with nonabsorb-
able sutures. Hormonal replacement was initiated 14 days after ovarioectomy by subcutaneous implantation of either 0.25 mg 17β-estradiol or placebo (vehicle) 21-day release pellets (Innovative Research of America). A cohort of these mice was immunized with PLP139–151, together with sham-ovariocto-
mized mice, to induce EAE 7 days after hormone replacement. Immunized animals were followed for EAE progression by monitoring clinical score and body weight. Tissues were obtained from sham-ovarioctomized and place-
bo-treated mice when they reached a score of 2 at 14 days after immuniza-
tion and from 17β-estradiol-treated mice, which remained free of disease, at 21 days after immunization. Protein lysates were prepared from CNS tis-
ues, while uteri and the upper third of vaginas of all mice were drop-fixed in 4% PFA for 48 hours. Following fixation, uterine tissue was trimmed of fat and connective tissue prior to determination of uterine weight.

EAE induction and in vivo treatment with JTE-013. EAE was induced in SJL/J mice via active immunization with proteolipid protein (PLP139–151; 200 μg) (GenScript USA Inc.) emulsified in complete Freund’s adjuvant containing Mycobacterium tuberculosis (H37Ra; Difco Laboratories). In addition, mice were injected with 200 ng pertussis toxin (List Biologicals Laboratories Inc.) at the time of immunization and 2 days after. Mice were graded for clinical manifestations of EAE by the following criteria: 1, tail weakness; 2, difficulty righting; 3, hind limb paralysis; 4, forelimb weakness or paralysis; 5, mori-
bund or dead. JTE-013 (1.5 mg/kg; Cayman Chemical Company) or vehicle (25% ETOH) was administered to mice daily via intraperitoneal injection, as previously described (46), when they achieved a clinical score of 2. B6129SF1 and Sipr2−/− mice were immunized similarly with 200 μg myelin oligodendro-
cyte glycoprotein (MOG35–55; GenScript USA Inc.) and 200 μg M. tuberculosis adjuvant. C57BL/6 mice were immunized as previously described (32).

In vivo assessment of BBB permeability. Mice were intraperitoneally injected with fluorescein sodium salt (100 mg/ml, Sigma-Aldrich), followed by collection of blood, intracardial perfusion, and harvesting of CNS tissues. Homogenates and sera were incubated overnight at 4°C in 1:1 dilution in 2% trichloroacetic acid (Sigma-Aldrich) and diluted in equal volumes of borate buffer, pH 11 (Sigma-Aldrich). Fluorescence emission at 538 nm was determined via a microplate reader, Synergy H1, and Gen5 software (both from BioTek Instruments Inc.). Tissue values were standardized against plasma values for individual mice.

Human subjects. The use of human postmortem biopsies for this study was approved by the Human Studies Committee of Washington University. Postmortem CNS tissue from 2 groups of patients was studied: 11 patients with clinically defined MS followed in the Washington University Multiple Sclerosis Center and 9 control individuals without histories of MS (Supplemental Table 2). The control group consisted of 8 patients without any clinical his-
tories of neurological disease and 1 patient with a history of CNS lymphoma.

IHC of S1PR2 in human CNS. Human postmortem specimens were col-
lected within 18 hours after death from 5 female and 5 male patients with MS and the same number of non-MS controls. Tissue preparation and evaluation of extent of inflammation were performed as previously described (31). For IHC, tissue sections were hydrated and fixed in 4% paraformaldehyde, blocked in 0.1% Triton X-100 and 10% goat serum, fol-
lowed by incubation with rabbit anti-S1PR2, rat anti-human CD31, mouse anti-human GFAP, and/or monoclonal mouse anti-human VE cadherin antibodies. Primary antibodies were detected with goat anti-rabbit Alexa Fluor 555, anti-rat Alexa Fluor 488, or goat anti-mouse Alexa Fluor 488 antibodies, as appropriate, followed by To-Pro-3 nuclear staining, confocal microscopy (Carl Zeiss USA), and quantitative analysis with Volocity 3D Image Analysis software (PerkinElmer Inc.). HRP-conjugated primary antibodies were detected via development in DAB solution (Cell Marque).

Histological and immunohistochemical analyses of murine samples. Murine CNS tissues underwent histological analyses (H&E; LFB staining) and IHC detection of CD31, GFAP, PDFG-Rβ, VE-cadherin, and CXCL12, as per-
formed previously (30). Analysis of polarized expression of CXCL12 across vasculature was performed as previously described (32). IHC detection of S1PR2 was performed in a fashion similar to that outlined for human CNS tissues. Visualization was done by confocal microscopy, as described above.

Preparation of in vitro human BBB. In vitro human BBB cultures were gen-
erated using HCMEC/D3, an endothelial cell line developed from brain tissue derived from the temporal lobe of an adult human female with epi-
lepsy (88). HCMEC/D3 cells were cocultured with primary human astro-
cytes (ScienCell), within a transwell system in which TEER values (Ω/cm²) were measured via chopstick electrode recording with an EVOM apparatus (World Precision Instruments), as previously described (47). Resistance val-
ues are reported as recorded values for each replicate minus the resistance of cell-free inserts (~90 Ω) measured alone. Measurements were taken 24 hours after initial seeding of endothelial cells (day 1); subsequent mea-
surements were taken daily for 10 days, at which point cultures were ready for experimentation. Changes in permeability were evaluated via TEER measurements after vehicle or S1P (Cayman Chemical Company) was added to top and bottom chambers (10–1,000 nM) and incubated for 2 hours, followed by addition of vehicle or JTE-013 (10–1,000 nM). Similar experi-
ments were done using the following agents: S1PR1-specific agonist, W146 (1,000 nM), and S1PR1-specific antagonist, SEW2871 (1,000 nM, both from Cayman Chemical Company), as well as caveolin endocytosis inhib-
itor, methyl-β-cyclodextrin (MBCD, 10 nM); clathrin endocytosis inhib-
itor, chlorpromazine (10 μg/ml); or the macropinocytosis/Pi3k inhibitor, wortmannin (100 nM, all from Sigma-Aldrich). To determine the effector molecules involved in S1PR2 signaling, we also pretreated for 2 hours with the Rac1 inhibitor, Z62954982 (1,000 nM, Cayman Chemical Company), the Rho/ROCK inhibitor, H-1152P (10 nM, Cayman Chemical Company), and the CDC42 inhibitor, ML141 (100 nM, Tocris, R&D Systems).

In vitro RNA interference. HCMEC/D3 cells in BBB cultures or grown on chamber slides were treated with 25 nM siRNA SMARTpools against S1PR1, S1PR2, or a nontargeting control pool (ON-TARGETplus, Thermo-
Scientific) complexed with 1:500 dilution of DharmaFect 1 transfection reagent (ThermoScientific) in antibiotic-free culture medium for 48 hours. Knockdown was confirmed via immunocytochemical analysis of S1PR1 and S1PR2 protein expression. Following siRNA treatment, cells were rinsed 3 times with PBS, returned to normal culture medium, and imme-
diately used for immunostaining or TEER measurement experiments.

Immunocytochemical evaluation of in vitro human BBB. Analysis of polarized expression of CXCL12 within HCMEC/D3 cells was performed after fixa-
tion with 4% paraformaldehyde for 60 minutes and permeabilization with 0.1% Triton X-100 and 10% goat serum for 30 minutes at room tempera-
ture. HCMEC/D3 cultures were incubated with primary antibodies against GGT-1, VE-cadherin, and biotinylated CXCL12 in 10% goat serum for 60 minutes at room temperature. Cells were washed 3 times in 1X dPBS and then incubated in goat anti-rabbit Alexa Fluor 555 and goat anti-
mouse Alexa Fluor 488 or Alexa Fluor 555 and streptavidin–Alexa Fluor 555 secondary antibodies in 10% goat serum for 15 minutes at room temperature. Cells were washed, counterstained with To-Pro-3, and cover-
slipped before being visualized via confocal microscopy.

Statistics. All microarray statistical analyses for the characterization of sexually dimorphic gene expression in female cerebellum were performed using Partek Genomics Suite (Partek Incorporated). Data were filtered for detectable probes (Illumina detection using P < 0.05 in at least one sample yielded 27,333 probes), quantile normalized by CNS region, and submitted to 2-way ANOVA for sex- and region-specific gene regulation. Transcripts that demonstrated upregulation or downregulation of >1.3 with a false discovery rate <0.2 were selected for further characterization. Two-tailed, Student’s t test was used to determine the statistical signif-
icance of immunohistochemical, mean maximal disease severity, and
cumulative clinical score analyses. Statistical significance of qRT-PCR, disease severity curve, fluorescent permeability, and TEER measurements was done by 2-way ANOVA, followed by Bonferroni’s post-hoc test with appropriate correction for repeated measures. Comparison of CXCL12 polarity was done via χ² test. For all graphs, error bars represent ± 1 SEM. All statistical analysis was performed with GraphPad Prism software, version 5.0. P values of less than 0.05 were considered significant for all comparisons.

Study approval. This study was carried out in strict accordance with the requirements pertaining to animal subjects within the Public Health Service Policy and USDA Animal Welfare Regulations. All experiments were performed in compliance with Washington University Institutional Animal Care and Use Committee (Animal Welfare Assurance A3381-01), which approved our animal protocol (20120160, 8/3/12). All efforts were made to minimize the suffering of animals used in this study. Postmortem human tissues banked for research purposes are exempt from institutional review board approval, and researchers do not have direct access to any identifying information.

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