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Blood-Brain Barrier Permeability Is Regulated by Lipid Transport-Dependent Suppression of Caveolae-Mediated Transcytosis

Highlights

- The unique lipid composition of CNS endothelial cells underlies BBB function
- Caveolae-mediated transcytosis is specifically suppressed downstream of Mfsd2a
- Lipids transported by Mfsd2a mediate the suppression of caveolae transcytosis
- Suppression of caveolae-mediated transcytosis is essential for BBB function

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In Brief

Andreone et al. revealed a key mechanism that explains how the suppression of transcytosis occurs to control BBB permeability. They discovered a single pathway revealing that the lipids transported by Mfsd2a control the formation of caveolae and therefore BBB integrity.



Neuron Article

Blood-Brain Barrier Permeability Is Regulated by Lipid Transport-Dependent Suppression of Caveolae-Mediated Transcytosis

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SUMMARY

The blood-brain barrier (BBB) provides a constant homeostatic brain environment that is essential for proper neural function. An unusually low rate of vesicular transport (transcytosis) has been identified as one of the two unique properties of CNS endothelial cells, relative to peripheral endothelial cells, that maintain the restrictive quality of the BBB. However, it is not known how this low rate of transcytosis is achieved. Here we provide a mechanism whereby the regulation of CNS endothelial cell lipid composition specifically inhibits the caveolae-mediated transcytotic route readily used in the periphery. An unbiased lipidomic analysis reveals significant differences in endothelial cell lipid signatures from the CNS and periphery, which underlie a suppression of caveolae vesicle formation and trafficking in brain endothelial cells. Furthermore, lipids transported by Mfsd2a establish a unique lipid environment that inhibits caveolae vesicle formation in CNS endothelial cells to suppress transcytosis and ensure BBB integrity.

INTRODUCTION

Vascular beds throughout the body have developed structural and functional heterogeneity to meet the demands of particular organs (Aird, 2007). The CNS requires a constant homeostatic microenvironment that is free of blood-borne toxins, allowing for proper synaptic transmission and information processing. The endothelial cells that line the walls of CNS blood vessels form a continuous, non-fenestrated vasculature. These cells constitute the blood-brain barrier (BBB), which regulates solute flux between blood circulation and the brain parenchyma (Andreone et al., 2015; Zhao et al., 2015). BBB disruption has been implicated in the pathogenesis of neurodegenerative diseases (Zhao et al., 2015; Zlokovic, 2008). Since BBB functionality is essential for normal brain activity, the impermeable nature of the BBB prevents brain uptake of most neurotherapeutic agents (Pardridge, 2012). This signifies a major obstacle to drug treatment of CNS diseases.

Pioneering electron microscopy (EM) work by Brightman, Reese, and Karnovsky (Brightman and Reese, 1969; Reese and Karnovsky, 1967) identified two structural features of CNS endothelial cells that underlie the restrictive properties of the BBB. First, specialized tight junctions prevent the paracellular passage of water-soluble molecules from the circulatory system to the brain parenchyma. Second, CNS endothelial cells exhibit unusually low levels of vesicle trafficking, a phenomenon that has generally been assumed to limit transcellular transport (transcytosis). While conventional wisdom has emphasized the importance of tight junctions, recent advances have highlighted the regulation of transcytosis as a crucial factor in limiting BBB permeability under physiological conditions and after pathological assaults (Armulik et al., 2010; Ben-Zvi et al., 2014; Daneman et al., 2010b; Knowland et al., 2014). However, the mechanism by which the suppression of transcytosis is achieved in CNS endothelial cells is completely unknown. An understanding of how transcytosis is regulated to maintain BBB integrity is not only fundamental to understanding BBB function, but also to developing therapeutic strategies to transiently open the BBB for CNS drug delivery.

Transcytotic pathways in epithelial and endothelial cells fall into two categories: clathrin mediated and non-clathrin mediated (Mayor et al., 2014). The mechanisms of clathrin-mediated vesicle trafficking are the most studied; these studies have identified the transferrin receptor-mediated transcytosis of iron as a key pathway that has been targeted for manipulation at the BBB for drug delivery (Bien-Ly et al., 2014; Niewoehner et al., 2014). Non-clathrin-mediated vesicle trafficking pathways utilize a broad range of large- and small-scale membrane invaginations that harness diverse cellular machinery, including dynamin, coat

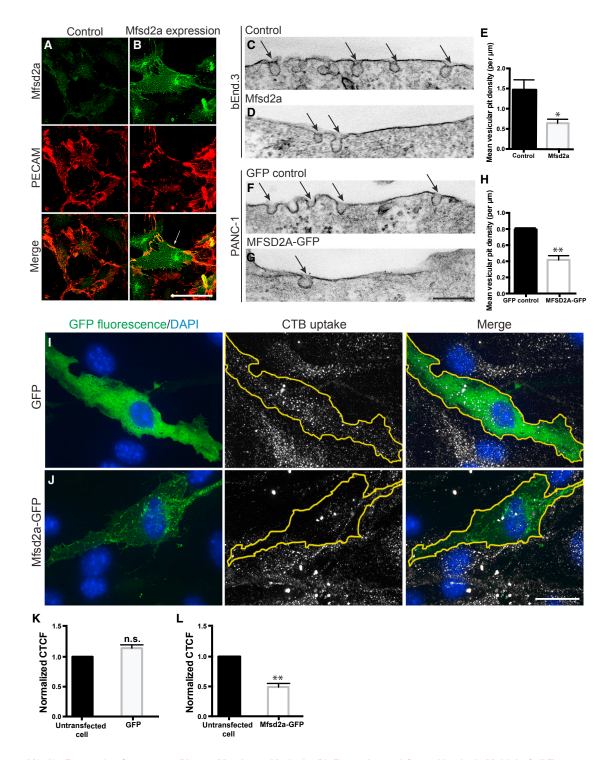


Figure 1. Mfsd2a Expression Suppresses Plasma Membrane Vesicular Pit Formation and Cargo Uptake in Multiple Cell Types

(A and B) Expression of Mfsd2a via lentivirus infection in bEnd.3 cells, in which endogenous Mfsd2a expression was not detectable (A). Mfsd2a (green) in bEnd.3 cells displays cell-surface localization (B) and co-localization with PECAM (red), a cell surface marker (arrows). Scale bar, 50 μ m.

(C–H) EM images showing the apical plasma membrane of bEnd.3 cells expressing mouse Mfsd2a (D) or PANC-1 cells expressing human MFSD2A-GFP (G). Mfsd2a-expressing cells show reduced vesicular pit density (arrows) in both bend.3 (E) and PANC-1 (H) cell types, compared to mock-infected control bEnd.3 cells (C) or PANC-1 cells expressing GFP alone (F). n = 3; 25 image profiles from at least 15 cells per experiment. Scale bar, 150 nm.

(I and J) Mfsd2a-expressing cells exhibit reduced cholera toxin subunit B (CTB) uptake. Uptake of CTB (gray; middle panels) in bEnd.3 cells expressing either soluble GFP (I) or Mfsd2a-GFP (J) (green, GFP fluorescence; blue, DAPI; left panels). Scale bar, 20 µm.

proteins, small GTPases, and RhoGAP proteins (Mayor et al., 2014). Among the many non-clathrin-mediated pathways, caveolae-mediated vesicle trafficking predominates as a transcytotic route used in the endothelial cells of peripheral organs, including heart and lung, to transport solutes from the blood to underlying tissue (Oh et al., 2007, 2014; Predescu et al., 1997; Schubert et al., 2001). Caveolae are 50-100 nm invaginations in the plasma membrane (Bastiani and Parton, 2010). The plasma membrane domains that form caveolae vesicles are highly ordered and enriched in saturated phospholipids, sphingolipids, ethanolamine plasmalogens (PE-p), and cholesterol (Parton and del Pozo, 2013; Pike et al., 2002). Additionally, the formation of caveolae vesicles requires the activity of both caveolin coat proteins and cytosolic adaptor proteins belonging to the cavin family (Bastiani and Parton, 2010). Although diverse functions have been identified for caveolae, their physiological role in different cell types and tissues is not fully understood (Cheng and Nichols, 2016).

Previously, we established that major facilitator superfamily domain containing 2a (Mfsd2a) is a key regulator for BBB function (Ben-Zvi et al., 2014). Mfsd2a is specifically expressed in CNS endothelial cells. Genetic ablation of Mfsd2a in mice results in a leaky BBB, as extravasation of several intravenously injected tracers, including 10 and 70 kD dextrans, NHS-biotin, and horseradish peroxidase (HRP), into the brain parenchyma is observed in $Mfsd2a^{-/-}$ mice from embryonic stages through adulthood. EM examination of brain endothelial cells from $Mfsd2a^{-/-}$ mice reveals increased transcytosis, as the intracellular vesicle number significantly increases. Additionally, HRP-injected mutants exhibit HRP-filled vesicles invaginating from the luminal plasma membrane, within the endothelial cell cytoplasm, and exocytosing at the abluminal plasma membrane. While these Mfsd2a^{-/-} mice have transcytotic defects, their tight junctions are normal. These findings demonstrate that Mfsd2a is specifically required to regulate transcytosis in CNS endothelial cells of the BBB and that the tracer extravasation observed in $Mfsd2a^{-/-}$ mice is solely due to increased levels of transcytotic vesicles within these cells. Therefore, Mfsd2a may serve as an ideal molecular handle to understand the mechanism by which transcytosis is regulated at the BBB.

There are several potential mechanisms whereby Mfsd2a regulates transcytosis in CNS endothelial cells, such as affecting transcytotic machinery via direct or indirect physical interactions or via controlling plasma membrane tension. Interestingly, in addition to its role in BBB function, Mfsd2a has been identified as a lipid transporter at the luminal plasma membrane of CNS endothelial cells to deliver the essential omega-3 fatty acid docosahexaenoic acid (DHA) into the brain (Nguyen et al., 2014). However, how DHA travels from the CNS endothelial cell plasma membrane to neurons is undetermined. DHA is heavily implicated in brain development (Innis, 2007), and, indeed, *Mfsd2a^{-/-}* mice have reduced total brain levels of DHA species and exhibit microcephaly (Nguyen et al., 2014). Therefore, although there is no existing example of lipids in endothelial cells playing a role in BBB function, examining if Mfsd2a's lipid transport function is related to BBB integrity will help narrow down the possible mechanisms of Mfsd2a-mediated regulation of transcytosis.

In this study, we show how transcytosis is suppressed in CNS endothelial cells to ensure proper BBB function by focusing on Mfsd2a. Using a combination of mouse genetics, lipidomic mass spectrometry, and EM analysis, we elucidate a pathway underlying the cellular mechanism of BBB function: lipids transported by Mfsd2a create a unique lipid composition of CNS endothelial cells that inhibits specifically caveolae-mediated transcytosis to maintain BBB integrity. Moreover, unbiased lipidomic analyses reveal that the lipid signatures of endothelial cells from the CNS and the periphery exhibit significant differences, which underlie the suppression of caveolae vesicle formation and trafficking in brain endothelial cells. Thus, this study establishes that lipid composition of CNS endothelial cells serves as a key new player in the regulation of transcytosis and barrier permeability.

RESULTS

Mfsd2a Cell-Autonomously Suppresses Vesicular Pit Formation and Cargo Uptake at the Plasma Membrane

To understand how transcytosis is regulated at the BBB, we first asked whether Mfsd2a is sufficient to suppress endocytic vesicle formation, the first step of all transcytotic pathways. Mfsd2a was expressed using lentivirus with high efficiency in the immortalized mouse brain endothelial cell line bEnd.3, in which endogenous Mfsd2a expression was not detectable (Figure 1A). Exogenous expression of Mfsd2a displayed cell-surface localization in bEnd.3 cells (Figure 1B), as well as cytoplasmic puncta, as seen in other cell lines (Nguyen et al., 2014; Reiling et al., 2011). EM revealed a reduction in the number of apical (analogous to luminal) micro-invaginations (pits) in cells expressing Mfsd2a compared to mock-infected control cells (Figures 1C-1E; Table S4A). A similar reduction in vesicular pit number was observed in a human pancreatic cancer cell line, PANC-1, stably transfected with human MFSD2A-GFP, compared to control cells stably transfected with GFP alone (Figures 1F-1H; Table S4A). These findings suggest that Mfsd2a expression is sufficient to inhibit vesicle formation.

To further assess whether Mfsd2a can directly inhibit cargo uptake, we next loaded confluent bEnd.3 cell cultures with a fluorescent cholera toxin subunit B (CTB) tracer (Figures 1I and 1J). Cells that express mouse Mfsd2a-GFP exhibited a 50% reduction in CTB uptake, as quantified by fluorescence intensity (Figure 1L), compared to cells that express GFP alone (Figure 1K). Use of the GFP-tagged constructs did not affect Mfsd2a protein localization (Reiling et al., 2011) (data not shown) and allowed for identification of positively transfected cells without quenching of the CTB fluorescence due to immunostaining protocol. These gain-of-function results are consistent with previous

⁽K and L) CTB uptake quantified as corrected total cell fluorescence (CTCF) per GFP transfected (K) or Mfsd2a-GFP transfected (L) cell compared to CTCF of at least two untransfected cells per image. n = 4; at least 15 transfected cells per experiment. All data are mean \pm SEM; n.s., not significant; *p < 0.05, **p < 0.01 (Student's t test). See also Table S4.

in vivo loss-of-function data (Ben-Zvi et al., 2014) and together demonstrate that Mfsd2a is necessary and sufficient for the suppression of vesicle formation and cargo uptake, cellular processes underlying transcytosis. Furthermore, they suggest that Mfsd2a acts via a direct mechanism to suppress CNS endothelial cell transcytosis in a cell-autonomous manner.

Mfsd2a Lipid Transport Function Is Required for Suppression of Transcytosis and BBB Integrity

To investigate how Mfsd2a may act directly at the CNS endothelial cell plasma membrane to regulate vesicle trafficking, we first asked whether the reported lipid transport function of Mfsd2a is required to suppress transcytosis. Although there is no known role of lipids in CNS endothelial cells for BBB function, answering this guestion will help to narrow down and pinpoint Mfsd2a's precise mechanism of action. Specifically to address this question, we generated a transporter-dead Mfsd2a knockin mouse as an in vivo tool. Asp96 is a conserved amino acid required for the cation binding that is crucial for transporter activity in MFS family members (Ethavathulla et al., 2014), and the D96A aspartic acid to alanine point mutation abolishes Mfsd2a transport function in vitro (Nguyen et al., 2014; Reiling et al., 2011). A mouse line harboring the Mfsd2a^{D96A} mutation in the endogenous Mfsd2a locus was generated via CRISPR/Cas9 gene editing. Specifically, one amino acid change resulting in the D96A missense mutation was introduced into exon 3 of Mfsd2a (Figures 2A, S1B, and S1C). Importantly, there was no observable difference in Mfsd2a expression pattern in Mfsd2a^{D96A/D96A} (hereafter denoted as Mfsd2a^{D96A}) mutants compared to Mfsd2a+/+ littermates by immunohistochemistry of neocortex and cerebellum (Figures 2B, 2C, S1D, and S1E). No difference in Mfsd2a expression was observed by western blot analysis of brain lysate (Figures 2F and 2G), indicating that Mfsd2a^{D96A} mutants express normal levels of Mfsd2a protein. Finally, Mfsd2a expression was restricted to BBB-containing CNS vasculature in Mfsd2a^{D96A} mutants, and was absent in non-BBB-containing vasculature of peripheral organs, including the lung and liver (Figures 2E and S1G), similar to Mfsd2a^{+/+} littermates (Figures 2D and S1F) (Ben-Zvi et al., 2014).

To examine the effect of the Mfsd2a^{D96A} mutation on the lipid transport function of Mfsd2a in vivo, we performed lipidomic mass spectrometry on brain lysate at postnatal stages. Lipidomic analysis of brain lysate revealed an altered lipid profile in Mfsd2a^{D96A} mutants (Figure S2A), with the level of a broad range of phospholipid species changed at postnatal day 4 (P4), including, but not limited to, DHA-containing species. Importantly, the lipid profile of Mfsd2a^{D96A} mutant brains closely mimicked that of $Mfsd2a^{-/-}$ mice, as the $Mfsd2a^{-/-}/Mfsd2a^{+/+}$ log ratio was very similar to the Mfsd2a^{D96A}/Mfsd2a^{+/+} log ratio of phospholipid levels measured by mass spectrometry (Figures 2H, S2A, and S2B; Tables S1 and S2). Together, these data demonstrate that the Mfsd2a^{D96A} mutation abolishes the lipid transport function of Mfsd2a in vivo without affecting protein expression, resulting in a lipid profile similar to that observed through global loss of Mfsd2a. Finally, Mfsd2a^{D96A} mutants exhibited microcephaly at P4 (Figure 2I), recapitulating the phenotype observed in *Mfsd2a^{-/-}* mice (Nguyen et al., 2014) and confirming that Mfsd2a lipid transport function is required for brain development.

control of BBB permeability. Previously, it was shown that Mfsd2a^{-/-} mice exhibit BBB leakage that is due to increased levels of transcytosis in CNS endothelial cells (Ben-Zvi et al., 2014). Because an increased vesicle number in CNS endothelial cells is the hallmark of the increased transcytosis phenotype observed in $Mfsd2a^{-/-}$ mice, we examined vesicle number in Mfsd2a^{D96A} mutants at P4 using EM (Figures 3A-3D). Mfsd2a^{D96A} mutants exhibited a significant increase in density of both luminal and abluminal membrane-connected vesicles (Figures 3E and 3F; Table S4B), similar to the phenotype observed in $Mfsd2a^{-/-}$ mice, indicating an upregulation of transcytosis. We next injected HRP into adult Mfsd2a^{D96A} mutants and Mfsd2a^{+/+} littermates and examined sub-cellular tracer localization. While HRP tracer was confined to the vessel lumen in Mfsd2a^{+/+} mice (Figures 3G and 3H), tracer in Mfsd2a^{D96A} mutants was observed being taken up into intracellular vesicles at the luminal plasma membrane, within the cytoplasm, and at the abluminal plasma membrane (Figures 3I and 3J), indicating that the upregulated vesicles in Mfsd2a^{D96A} mutants act as transcytotic carriers. In both Mfsd2a+/+ wild-type littermates and Mfsd2a^{D96A} mice, HRP tracer entered the space between CNS endothelial cells but was sharply halted at tight junction "kissing points" (Figures 3K and 3L), indicating that tight junctions are functional. Together, these data demonstrate that loss of Msfd2a transport function results in increased CNS endothelial cell transcytosis, without impacting tight junction integrity.

With this new genetic tool, we next asked if Mfsd2a lipid trans-

port function is necessary for suppression of transcytosis and

To determine whether the observed increased transcytosis results in BBB leakage in Mfsd2a^{D96A} mutants, we performed intravenous injection of fluorescent tracer at P4 and examined tracer localization as a metric of BBB integrity. We chose 10 kD dextran tracer for analysis, as robust extravasation of this tracer was previously observed in $Mfsd2a^{-/-}$ mice (Ben-Zvi et al., 2014). While tracer was completely confined within the vasculature of $Mfsd2a^{+/+}$ littermate control animals (Figure 3M). tracer extravasation was observed throughout the brain of Mfsd2a^{D96A} mutants (Figure 3N), with tracer diffused into the brain parenchyma and taken up by non-vascular cells. Permeability index quantification (defined as area of tracer/area of vessels) revealed a statistically significant increase of BBB leakage in Mfsd2a^{D96A} mutants compared to Mfsd2a+/+ controls (Figure 3O). BBB leakage in Mfsd2a^{D96A} mutants persisted into adulthood, as HRP tracer extravasation into the forebrain parenchyma was observed at P90 (Figures S3A and S3B). Taken together, these data demonstrate that BBB leakage occurs in Mfsd2a^{D96A} mice via increased transcytosis, similar to $Mfsd2a^{-/-}$ mice. Furthermore, they strongly implicate the lipid transport function of Mfsd2a in the suppression of vesicle trafficking.

Brain Capillaries Possess a Lipid Signature Underlying Suppression of Caveolae

Our results demonstrate that Mfsd2a cell-autonomously suppresses vesicle trafficking via a lipid transport-dependent mechanism. This raises the possibility that lipid transport directly affects the membrane lipid composition of CNS endothelial cells, which in turn may result in the suppression of transcytosis at the BBB. To further understand the link between lipids and vesicle

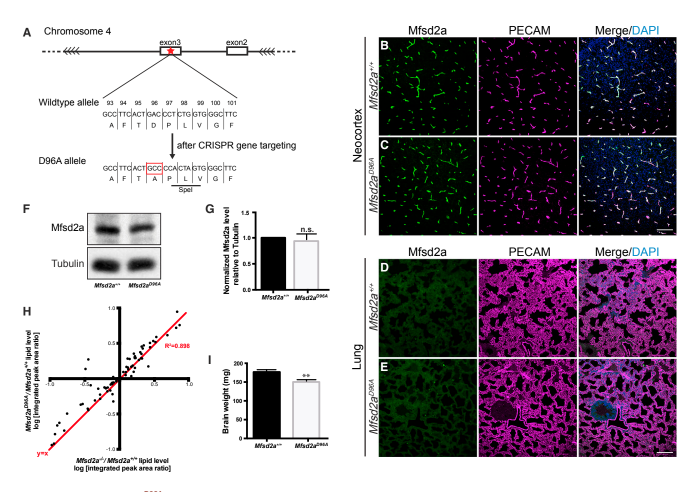


Figure 2. A Novel Mfsd2a^{D96A} Mouse Abolishes the Lipid Transport Function of Mfsd2a and Displays Microcephaly

(A) Gene targeting strategy for the generation of *Mfsd2a*^{D96A} mice. Magnified view of the genomic region that contains residue D96 in exon 3 of *Mfsd2a*. CRISPR/ Cas9 genome editing introduces the D96A point mutation (red box), as well as two additional silent mutations in codons 97/98 that generate an Spel restriction site (black underline) for screening purposes. For additional details, see <u>STAR Methods</u>.

(B and C) Immunostaining at P4 of Mfsd2a protein (green) reveals vascular expression (purple, PECAM) in the neocortex of both $Mfsd2a^{+/+}$ control (B) and $Mfsd2a^{D96A}$ mutants (C). n = 3 animals per genotype. Scale bar, 100 μ m.

(D and E) Immunostaining at P4 reveals absence of Mfsd2a protein expression in non-BBB-containing lung vasculature (purple, PECAM) of both *Mfsd2a*^{+/+} control (D) and *Mfsd2a*^{D96A} mutants (E). n = 3 animals per genotype. Scale bar, 100 μm.

(F) Western blot from P4 brain lysate of Mfsd2a^{D96A} mutants and littermate Mfsd2a^{+/+} controls. Tubulin (loading control).

(G) Quantification shows equal Mfsd2a protein levels between $Mfsd2a^{D96A}$ mutants and littermate $Mfsd2a^{+/+}$ controls. n = 3 littermate pairs per genotype. Data are mean \pm SEM; n.s., not significant (Student's t test).

(H) The lipid profile of $Mfsd2a^{D96A}$ mutant brains closely mimics that of $Mfsd2a^{-/-}$ mutant brains at P4. Dot plot representation of $Mfsd2a^{-/-}/Mfsd2a^{+/+}$ log ratio (x axis), compared to $Mfsd2a^{D96A}/Mfsd2a^{+/+}$ log ratio (y axis) of phospholipid levels measured by mass spectrometry. Each dot represents a single lipid species; red line, y = x; R² = 0.898. $Mfsd2a^{D96A}$ experiment, n = 6 animals per genotype; $Mfsd2a^{-/-}$ experiment, n = 4 animals per genotype.

(I) *Mfsd2a*^{D96A} mutants exhibit microcephaly. Quantification of brain weight at P4. n = 5 animals per genotype. Data are mean ± SEM; **p < 0.01 (Student's t test). See also Figures S1 and S2 and Tables S1 and S2.

trafficking, we next asked whether the lipid composition of CNS endothelial cells, a previously unexplored facet of BBB function, shows any differences compared to peripheral endothelial cells. To investigate this possibility, we performed lipidomic mass spectrometry analysis on acutely isolated capillaries from brain and lung of wild-type mice at P4. Lung vasculature does not possess the restrictive barrier properties of CNS vasculature, and lung endothelial cells are enriched in transcytotic vesicles, which have been shown to transport cargo from the circulation to the lung parenchyma (Ghitescu et al., 1986; Oh et al., 2007; Schubert et al., 2001). Western blot analysis showed that this capillary preparation is highly enriched in endothelial cells in both brain and lung, indicated by an abundance of VE-cadherin protein compared to total tissue lysate (Figure S4B). Residual pericytes were present in both brain and lung capillary preparations, and, importantly, the brain capillary preparation was free of both neurons (NeuN) and astrocyte endfeet (Aquaporin-4) (Figures S4A and S4B).

Lipidomic analysis revealed that although most lipid species were found at similar levels in isolated brain and lung capillaries, significant differences in several lipid species were observed (Figures 4A and S4C; Table S3). Remarkably, one of the most

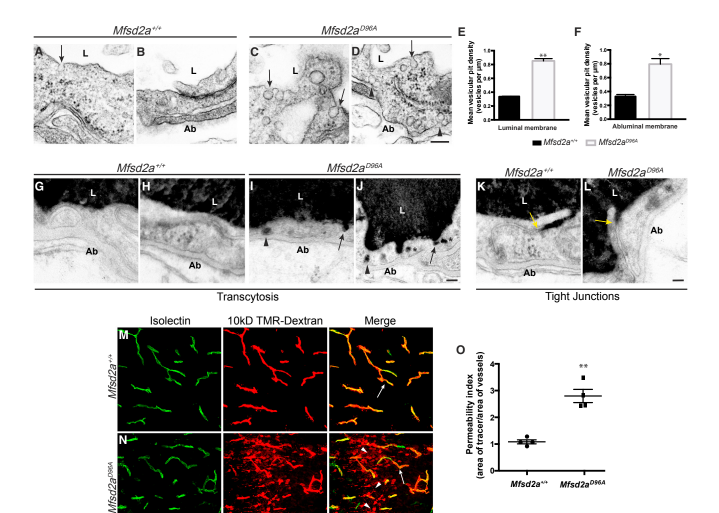


Figure 3. The Lipid Transport Function of Mfsd2a Is Required for the Suppression of Transcytosis and Maintenance of BBB Integrity In Vivo (A–D) Increased vesicular density in CNS endothelial cells of *Mfsd2a*^{D96A} mutants (C and D) compared to their littermate controls (A and B). EM analysis at P4 reveals an increased number of vesicles in *Mfsd2a*^{D96A} mutants along the luminal (arrows) and abluminal (Ab; black arrowheads) plasma membranes. Scale bar, 150 nm.

(E and F) Quantification of vesicular density along the luminal (E) and abluminal (F) plasma membranes. n = 3 animals per condition; 20 capillary cross-sections per animal.

(G–J) Increased transcytosis is evident in CNS endothelial cells of HRP-injected $Mfsd2a^{D96A}$ mutants at P90. HRP tracer (black) is confined within the vessel lumen (L) of littermate control animals (G and H). Tracer-filled vesicles are observed invaginating from the luminal plasma membrane (arrows), within the cytoplasm (asterisks), and along the abluminal plasma membrane (arrowheads) in $Mfsd2a^{D96A}$ mutants (I and J). n = 3 animals per condition. Scale bar, 100 nm.

(K and L) Tight junctions are functional in CNS endothelial cells of HRP-injected $Mfsd2a^{D96A}$ mutants. HRP tracer fills the vessel lumen (L) and enters intercellular clefts but is sharply halted at tight junction "kissing points" (yellow arrows) in both $Mfsd2a^{D96A}$ mutants (L) and $Mfsd2a^{+/+}$ littermate control animals (K) at P90. n = 3 animals per condition. Scale bar, 100 nm.

(M and N) *Mfsd2a^{D96A}* mutants display BBB leakage. 10 kD TMR-dextran tracer (red) is completely confined within vessels (green, Isolectin B4 [IB4]) (arrows) in *Mfsd2a^{+/+}* littermate control animals (M) at P4. Tracer-filled parenchyma cells (arrowheads) surround vessels in the brain of *Mfsd2a^{D96A}* mutants (N). Scale bar, 50 μm.

(O) Permeability index of tracer leakage in brain, as quantified by area of tracer divided by area of vessels per image (value = 1 signifying no leakage). n = 4 animals per genotype. All data are mean ± SEM; *p < 0.05, **p < 0.01 (Student's t test). See also Figures S3 and S6 and Table S4.

significantly enriched lipid populations in brain capillaries was DHA-containing phospholipid species in lysophosphatidylethanolamine (LPE), phosphatidylcholine (PC), phosphatidylethanolamine (PE), and phosphatidylserine (PS) subclasses, ranging from 1.9- to 5.1-fold enrichment (Figure 4B). Significant enrichment of a few non-DHA-containing lipids, such as C34:0 PC and C38:4 PE, was also observed in brain capillaries (Figure 4A). On the other hand, modest decreases in several PE-p species

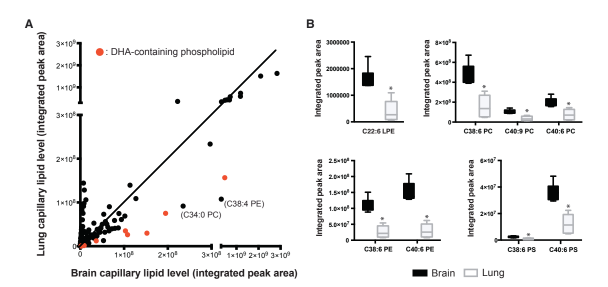


Figure 4. Lipidomic Profiling of Brain versus Lung Capillaries Reveals Different Lipid Signatures, Including Changes in DHA-Containing Phospholipid Species

(A) Dot plot representation of lipidomic profiling of brain versus lung capillaries from P4 wild-type mice. Dots represent average lipid level in the brain (x axis) and lung (y axis) for each metabolite tested, expressed as mass spectrometry integrated peak area. Red dots indicate DHA-containing phospholipid species.
(B) DHA-containing phospholipid levels are increased in brain capillaries. Lipidomic analysis of DHA-containing phospholipid species from brain and lung capillaries of P4 wild-type mice. Value for each species is normalized to internal standard and expressed as integrated mass spectrometry peak area (LPE, lyso-phosphatidylethanolamine; PC, phosphatidylcholine; PE, phosphatidylethanolamine; PS, phosphatidylserine). *p < 0.004 (Student's t test, with Bonferroni correction for multiple comparisons). For all data, n = 6 groups per organ; 8 animals pooled per group.
See also Figures S4 and S5 and Table S3.

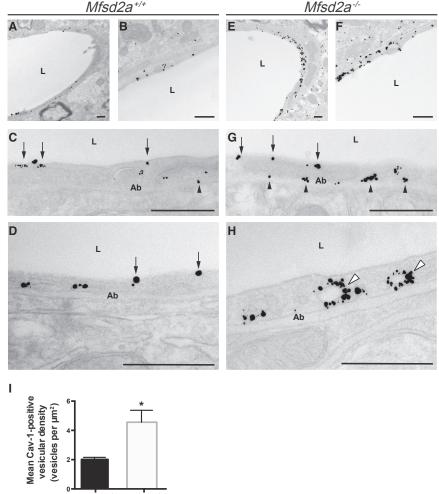
were observed in brain capillaries compared to lung capillaries (Figure S5A). Although both the acutely isolated brain and lung capillary preparations retained residual pericytes, it unlikely that these cells contribute to the observed differences in lipid signature, for pericytes do not express Mfsd2a (Ben-Zvi et al., 2014), the only known DHA transporter in the brain; there is no evidence that pericytes are able to synthesize DHA de novo; and both brain and lung capillary preparations were highly enriched in endothelial cells.

Notably, lung endothelial cells have abundant caveolae vesicles that can act as transcytotic carriers (Ghitescu et al., 1986; Oh et al., 2007; Schubert et al., 2001). The formation and functionality of caveolae membrane domains can be affected by plasma membrane lipid composition (Lingwood and Simons, 2010; Parton and del Pozo, 2013). In particular, DHA is a highly unsaturated fatty acid that disrupts liquid-ordered caveolae domains, normally enriched in saturated phospholipids, sphingolipids, PE-p species, and cholesterol. DHA displaces Caveolin-1 (Cav-1), a coat protein of the caveolin family that is obligatory for caveolae vesicle formation, from the plasma membrane. Thus, enrichment of DHA in the plasma membrane renders caveolae membrane domains less capable of forming caveolae vesicles (Chen et al., 2007; Li et al., 2007; Ma et al., 2004). Therefore, the lipid composition of brain capillaries enriched in DHA species and low in PE-p species may reflect a plasma membrane microenvironment in CNS endothelial cells that is unfavorable for the assembly of functional caveolae domains and subsequent vesicle formation. Together, these analyses reveal significant differences in the lipid signature of BBB-containing brain capillaries versus non-BBB-containing lung capillaries, some of which may contribute to the suppression of transcytosis at the BBB by inhibiting caveolae vesicle formation and trafficking in CNS endothelial cells.

Significantly reduced levels of several triglycerides (TAGs) and cholesteryl ester (CE) species were observed in brain capillaries versus lung capillaries (Figures S5B and S5C). TAGs and CEs are stored in intracellular lipid droplets (Guo et al., 2009), and both in vitro and in vivo evidence suggests that the storage of these lipids depends on caveolae vesicle trafficking (Cohen et al., 2004; Martin and Parton, 2005; Pol et al., 2004). Therefore, low levels of TAGs and CEs in brain capillaries may reflect a low level of caveolae trafficking. Together these observations raise the possibility that caveolae vesicles may be the subtype increased in CNS endothelial cells that accounts for the leaky BBB in $Mfsd2a^{-/-}$ mice.

Caveolae Vesicle Trafficking Is Upregulated in *Mfsd2a^{-/-}* Mice

To test this hypothesis, we performed Cav-1 immuno-EM labeling in brains of $Mfsd2a^{-/-}$ mice and control littermates (Figures 5A-5H). Cav-1 immunoreactivity (gold particles) was seen along the luminal plasma membrane in CNS endothelial cells of $Mfsd2a^{+/+}$ control mice, whereas immunoreactivity in $Mfsd2a^{-/-}$ mice was observed along the luminal and abluminal plasma membranes (Figure 5G), and surrounding vesicle structures in the cytoplasm (Figure 5H). Additionally, there was a significant increase in the number of Cav-1-positive vesicles in CNS endothelial cells of $Mfsd2a^{-/-}$ mice compared to controls



Mfsd2a⁺/⁺ Mfsd2a⁻∕-

(Figure 5I; Table S4C). These data suggest that caveolae vesicle trafficking is upregulated in mice lacking *Mfsd2a*, and that Mfsd2a may normally act to suppress caveolae-mediated transcytosis in CNS endothelial cells.

Caveolae-Mediated Transcytosis Is Specifically Suppressed Downstream of Mfsd2a Function at the BBB In Vivo

To directly test the idea that the caveolae-mediated transcytosis pathway is suppressed downstream of Mfsd2a function at the BBB, we performed a genetic rescue experiment by crossing $Mfsd2a^{-/-}$ with $Cav-1^{-/-}$ mice, as Cav-1 is obligatory for caveolae formation. While $Cav-1^{-/-}$ mice are viable, they are unable to form caveolae vesicle structures and display defects in cargo uptake in vivo (Razani et al., 2001; Schubert et al., 2001). If the caveolae pathway is downstream of Mfsd2a in vivo, then crossing $Cav-1^{-/-}$ mice with $Mfsd2a^{-/-}$ mice should reverse the upregulated transcytosis seen in $Mfsd2a^{-/-}$ mice, as CNS endothelial cells of $Mfsd2a^{-/-}$; $Cav-1^{-/-}$ double knockouts will no longer be able to form caveolae vesicles to act as transcytotic carriers. Alternatively, if Mfsd2a suppresses other transcytotic

(A–H) Cav-1 immunoreactivity in CNS endothelial cells of $Mfsd2a^{+/+}$ and $Mfsd2a^{-/-}$ mice under EM. EM analysis with immuno-gold labeling of Cav-1 protein in CNS endothelial cells of $Mfsd2a^{+/+}$ (A–D) and $Mfsd2a^{-/-}$ mice (E–H) at P90. Gold particles (black puncta) are visualized along the luminal (arrows) and abluminal (Ab; black arrowheads) plasma membranes, as well as within the cytoplasmic particles are seen associated with vesicular structures (white arrowheads). All scale bars, 500 nm.

(I) Quantification of Cav-1-positive vesicles in $Mfsd2a^{+/+}$ and $Mfsd2a^{-/-}$ mice. n = 3 animals per genotype; 10 capillary cross-sections per animal. Data are mean \pm SEM; *p < 0.05 (Student's t test). See also Table S4.

pathways, increased transcytosis will still be observed in *Mfsd2a^{-/-}; Cav-1^{-/-}* double knockouts.

Similar to previous results (Ben-Zvi et al., 2014), EM analysis revealed occasional vesicles in $Mfsd2a^{+/+}$; $Cav-1^{+/+}$ wild-type CNS endothelial cells at P5 (Figures 6A and 6B), with an over 2.5-fold increase in total luminal and abluminal membrane-connected vesicle density in $Mfsd2a^{-/-}$; $Cav-1^{+/+}$ single knockouts (Figures 6C, 6D, 6I, and 6J). Consistent with the finding that $Cav-1^{-/-}$ mice are unable to form caveolae vesicles, we observed a nearly 2-fold decrease in vesicle density in $Mfsd2a^{+/+}$; $Cav-1^{-/-}$ single knockouts compared to wild-type

levels (Figures 6E, 6F, 6I, and 6J). However, in *Mfsd2a^{-/-}; Cav*-1^{-/-} double knockouts, a complete reversal of the *Mfsd2a^{-/-}* single mutant phenotype was evident, with vesicle density mimicking that of *Cav*-1^{-/-} single mutants (Figures 6G–6J; Table S4D). Therefore, the caveolae-mediated transcytosis pathway is the one that is normally suppressed by Mfsd2a. In contrast, no significant difference in the density of clathrin-coated pits, identified under EM by the presence of an electron-dense clathrin coat (Figure S6A), was observed in *Mfsd2a^{-/-}, Cav*-1^{-/-}, *Mfsd2a^{-/-}; Cav*-1^{-/-}, or *Mfsd2a^{D96A}* mutants compared to their littermate controls (Figures S6B–S6E; Tables S4E and S4F). Together, these data demonstrate that Mfsd2a acts upstream of Cav-1 specifically to suppress the caveolae vesicle trafficking pathway at the BBB.

Finally, we asked if caveolae-mediated transcytosis is required for regulating BBB permeability by examining whether genetic loss of *Cav-1* is able to rescue the BBB leakage observed in the brains of *Mfsd2a^{-/-}* mice. To evaluate BBB permeability, we again examined 10 kD dextran tracer localization in the brain after intravenous injection at P5. Tracer was confined within the brain vasculature of *Mfsd2a^{+/+}; Cav-1^{+/+}* wild-type and

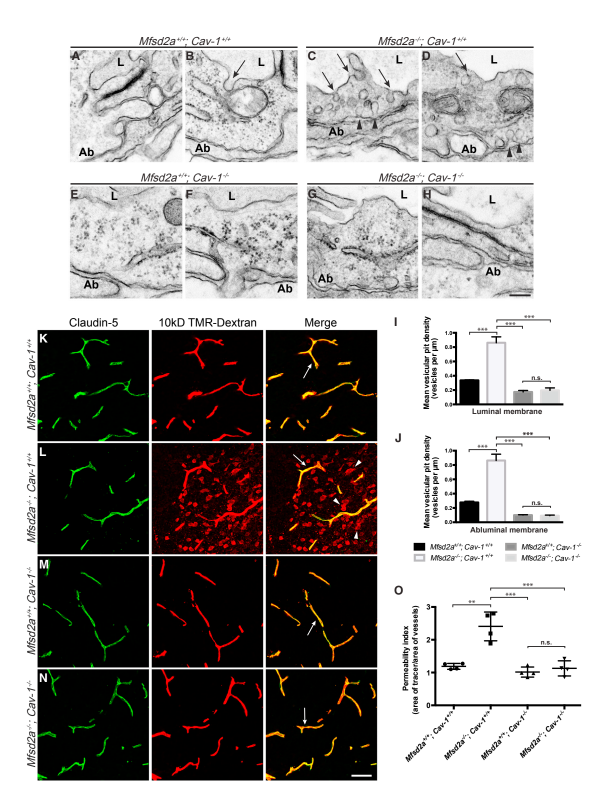
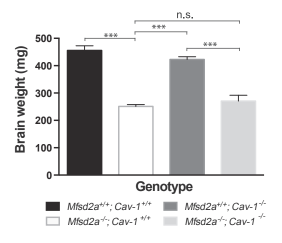


Figure 6. Genetic Inhibition of the Caveolae Pathway Rescues Both the Increased Vesicular Density and BBB Leakage Phenotypes of *Mfsd2a^{-/-}* Mice

(A–H) Genetic inhibition of the caveolae pathway rescues increased vesicular density in $Mfsd2a^{-/-}$ animals. EM examination of BBB integrity in $Mfsd2a^{+/+}$; $Cav-1^{+/+}$ (A and B), $Mfsd2a^{-/-}$; $Cav-1^{+/+}$ (C and D), $Mfsd2a^{+/+}$; $Cav-1^{-/-}$ (E and F), and $Mfsd2a^{-/-}$; $Cav-1^{-/-}$ (G and H) mice at P5. Increased vesicular pit density in $Mfsd2a^{-/-}$; $Cav-1^{+/+}$ single mutant CNS endothelial cells is observed along the luminal (arrows) and abluminal (Ab; arrowheads) plasma membranes. $Mfsd2a^{+/+}$; $Cav-1^{+/+}$ wild-type, $Mfsd2a^{+/+}$; $Cav-1^{-/-}$ ingle mutant, and $Mfsd2a^{-/-}$; $Cav-1^{-/-}$ double mutant cells exhibited very few vesicles. Scale bar, 150 nm.

(legend continued on next page)





Quantification of brain weight at P90. n = 5 animals per genotype. Data are mean \pm SEM; n.s., not significant; ***p < 0.001 (one-way ANOVA, post hoc Tukey test).

Mfsd2a^{+/+}; Cav-1^{-/-} single knockouts, indicating no BBB leakage in these mice (Figures 6K and 6M). Tracer extravasation was observed in Mfsd2a^{-/-}; Cav-1^{+/+} single knockouts, as tracer diffused into the brain and was taken up by non-vascular parenchymal cells (Figure 6L). A significant increase in the permeability index quantification of Mfsd2a^{-/-}; Cav-1^{+/+} single knockouts was observed, compared to both Mfsd2a+/+; Cav-1+/+ wildtype and *Mfsd2a*^{+/+}; *Cav-1*^{-/-} single knockouts (Figure 6O). However, no leakage was observed in Mfsd2a^{-/-}; Cav-1^{-/-} double knockouts, as tracer was confined within the brain vasculature and the permeability index quantification revealed no significant difference to that of both Mfsd2a+/+; Cav-1+/+ wild-type and *Mfsd2a*^{+/+}; *Cav*-1^{-/-} single knockouts (Figures 6N and 6O). This observed rescue of BBB leakage was also evidenced at adult stages, as extravasation of HRP tracer into the forebrain parenchyma was observed in Mfsd2a^{-/-}; Cav-1^{+/+} single knockouts (Figure S3D), but not wild-type, $Cav-1^{-/-}$ single knockout, or Mfsd2a^{-/-}; Cav-1^{-/-} double knockout mice at P90 (Figures S3C, S3E, and S3F). Consistent with previous findings (Knowland et al., 2014), we found that tight junctions were functional in Cav-1^{-/-} mice, as HRP tracer was halted at tight junction "kissing points" in both Cav-1 mutants (Figures S7C and S7D) and wild-type littermates (Figures S7A and S7B) under EM. Together with our vesicle density analysis, these data indicate that increased caveolae vesicle trafficking results in the BBB leakage observed in Mfsd2a^{-/-} mice, as genetic loss of Cav-1

was able to rescue tracer extravasation. Furthermore, of the numerous vesicular trafficking pathways currently known, these data demonstrate that Mfsd2a maintains BBB integrity by specifically suppressing the caveolae-mediated transcytotic pathway.

Caveolae-Mediated Transcytosis Is Not Required for Delivery of DHA into the Brain Parenchyma for Brain Development

Interestingly, although $Mfsd2a^{-/-}$; $Cav-1^{-/-}$ double knockouts were able to rescue the BBB defects seen in $Mfsd2a^{-/-}$ mice, they were unable to rescue the microcephaly phenotype (Figure 7). Thus, the caveolae pathway downstream of Mfsd2a in CNS endothelial cells specifically controls transcytosis and BBB function, but is not required for Mfsd2a-mediated delivery of DHA into the brain parenchyma for proper brain development. Our analysis of the $Mfsd2a^{D96A}$ mouse, together with the comparative brain and lung capillary lipidomic profiling, indicates that the suppression of the caveolae pathway requires the transport of lipids, notably DHA-containing phospholipids, by Mfsd2a to regulate CNS endothelial cell plasma membrane composition and inhibit caveolae vesicle formation.

DISCUSSION

While it was observed nearly 50 years ago that CNS endothelial cells exhibit unusually low levels of transcytotic vesicles (Brightman and Reese, 1969; Reese and Karnovsky, 1967), it has remained poorly understood whether this low rate of transcytosis is important for BBB function, what type of vesicles are involved, and how vesicle formation and transcvtosis are maintained at low levels in brain endothelial cells. In this study, we reveal that the unique membrane lipid composition of CNS endothelial cells regulates BBB permeability by suppressing caveolae-mediated transcytosis. Our findings support a model in which Mfsd2amediated transport of DHA controls the luminal plasma membrane DHA lipid composition of CNS endothelial cells such that it actively prevents the formation of functional caveolae membrane domains (Figures 8A and 8B). Thus, unlike lung endothelial cells, which have low levels of DHA and display high levels of transcytotic caveolae vesicles, Mfsd2a-mediated lipid transport into CNS endothelial cells renders them less capable of forming caveolae vesicles to act as transcytotic carriers. As a result, the BBB is functional. Together, these findings uncover a mechanism for the suppression of transcytosis in CNS endothelial cells to maintain BBB integrity, and reveal an essential role for the precise control of membrane lipid composition in the regulation of BBB permeability.

(O) Permeability index of tracer leakage in brain, as quantified by area of tracer divided by area of vessels per image (value = 1 signifying no leakage). n = 4 animals per genotype. All data are mean \pm SEM; n.s., not significant; **p < 0.01, ***p < 0.001 (one-way ANOVA, post hoc Tukey test). See also Figures S3, S6, and S7 and Table S4.

⁽I and J) Quantification of vesicular pit density at the luminal (I) and abluminal (J) plasma membranes. n = 4 animals per genotype; 20 capillary cross-sections per animal.

⁽K–N) Genetic inhibition of the caveolae pathway rescues BBB leakage defects in $Mfsd2a^{-/-}$ animals. 10 kD dextran tracer (red) was injected into the circulation of $Mfsd2a^{+/+}$; $Cav-1^{+/+}$ (K), $Mfsd2a^{-/-}$; $Cav-1^{+/+}$; $Cav-1^{+/+}$ (K), $Mfsd2a^{-/-}$; $Cav-1^{+/+}$ (K), Mfsd2a

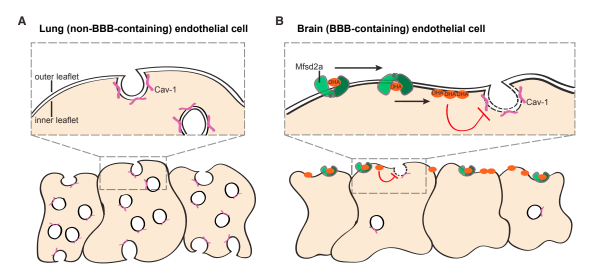


Figure 8. Model for the Suppression of Caveolae-Mediated Transcytosis via Regulated Membrane Lipid Composition at the BBB (A) Non-BBB-containing lung endothelial cells do not express Mfsd2a, display low levels of DHA, and display high levels of transcytotic caveolae vesicles, characterized by the presence of Cav-1 coat protein (purple).

(B) BBB-containing brain endothelial cells express Mfsd2a (green). Mfsd2a acts as a lipid flippase, transporting phospholipids, including DHA-containing species (orange), from the outer to inner plasma membrane leaflet. The increased levels of DHA, and presumably other lipid changes, alter the plasma membrane composition such that caveolae vesicles are unable to form and act as transcytotic carriers. Thus, brain endothelial cells display low levels of caveolae vesicles. This suppression of caveolae formation and trafficking subsequently ensures BBB integrity under normal conditions.

The Unique Lipid Composition of CNS Endothelial Cells Underlies BBB Function

Although the field has successfully identified several key gene and protein regulators for BBB function (Ben-Zvi et al., 2014; Chow and Gu, 2015; Daneman et al., 2010a; Junge et al., 2009; Liebner et al., 2008; Sohet et al., 2015; Tam et al., 2012; Zhou et al., 2014), the question of whether lipids can regulate BBB function has yet to be examined. Our demonstration of the in vivo requirement of the lipid transport function of Mfsd2a for the suppression of transcytosis and BBB permeability, coupled with our result of lipidomic profiling of capillaries from brain versus lung, reveals lipid composition of CNS endothelial cells as a key regulator for BBB function. Previous work on Mfsd2a has focused on the deleterious consequences of decreased DHA levels in brain parenchymal cells, such as behavioral abnormalities and microcephaly (Alakbarzade et al., 2015; Guemez-Gamboa et al., 2015; Nguyen et al., 2014). Our results demonstrate an additional direct functional role of DHA transport at the CNS endothelial cell luminal plasma membrane, the site of Mfsd2a expression in the brain, for suppression of caveolae transcytosis and maintenance of BBB integrity. Besides DHA species, C34:0 PC and C38:4 PE are also significantly enriched in brain capillaries relative to lung capillaries, and their function at the BBB remains to be determined. Together, our Mfsd2a^{D96A} mouse model and brain versus lung lipidomic profiling illuminate a previously unknown role of CNS endothelial cell lipid composition in BBB integrity. These findings may usher in new lines of research into how the metabolomic, in addition to the proteomic, signature of CNS endothelial cells regulates BBB function.

Suppression of Caveolae-Mediated Transcytosis Is Essential for BBB Function

Compared to the relatively well-studied mechanisms of clathrinmediated vesicle trafficking, the physiology of caveolae is less understood. Caveolae have multiple putative functions at the plasma membrane, including cell signal transduction, mechanosensation, lipid metabolism, and plasma membrane organization, and their physiological role in different cell types and tissues is not fully clear (Cheng and Nichols, 2016; Parton and del Pozo, 2013). While endothelial cells in peripheral organs, such as the lung and heart, are enriched in caveolae, and previous studies have demonstrated that caveolae vesicles act as transcytotic carriers in these peripheral organ endothelial cells (Ghitescu et al., 1986; Oh et al., 2007, 2014; Schubert et al., 2001), there remains a long-standing debate as to what role, if any, caveolae play in transcytosis. In particular, recent in vitro studies have challenged the notion that caveolae significantly contribute to endocytic or transcytotic flux (Bitsikas et al., 2014; Shvets et al., 2015). Our genetic rescue experiments demonstrate that the presence of transcytotic caveolae vesicles accounts for BBB leakage in the absence of Mfsd2a, and thus assign a physiological role to caveolae-mediated transcytosis in CNS endothelial cells to regulate BBB permeability.

Regulated Lipid Composition Suppresses Caveolae-Mediated Transcytosis for BBB Function

The mutation of a single amino acid caused $Mfsd2a^{D96A}$ transporter-dead mice to phenocopy the increased transcytosis and BBB leakage defects observed in $Mfsd2a^{-/-}$ global knockout mice. Our unbiased lipidomic analysis showing DHA species as one of the most significantly enriched lipid populations in brain

capillaries compared to lung capillaries further points to a direct role of DHA in the suppression of caveolae vesicles. How does DHA inhibit caveolae vesicles? Caveolae formation depends on both Cav-1 and cholesterol (Lingwood and Simons, 2010; Parton and del Pozo, 2013). Our findings support previous studies that have shown that DHA treatment of human retinal endothelial cells and human umbilical vein endothelial cells resulted in a significant depletion of cholesterol in the membrane as well as the displacement of Cav-1 and several signaling proteins resident in caveolae (Chen et al., 2007; Li et al., 2007). Moreover, in vivo studies show that feeding mice with omega-3 fatty acid-enriched diet resulted in a concomitant reduction of cholesterol and Cav-1 in mouse colon epithelium (Ma et al., 2004). Therefore, it is likely that in CNS endothelial cells, Mfsd2a acts as a lipid flippase (Quek et al., 2016), transporting phospholipids, including DHA-containing species, from the outer to the inner plasma membrane leaflet. The increased levels of DHA in the inner plasma membrane leaflet subsequently cause the displacement of cholesterol and Cav-1, thereby inhibiting caveolae formation. It is plausible that DHA enrichment in the plasma membranes results in higher membrane fluidity and low membrane curvature that is not favorable for caveolae formation.

In light of the emerging evidence in both the normal and diseased brain that transcytosis represents a major mechanism for BBB function (Armulik et al., 2010; Ben-Zvi et al., 2014; Chow and Gu, 2017; Daneman et al., 2010b; Haley and Lawrence, 2017; Knowland et al., 2014), our work provides a mechanistic understanding of how transcytotic pathways are suppressed in the CNS to ensure this function. Future studies, including examining the effect of acute removal of Mfsd2a function in adult mice and the effect of dietary manipulation of lipids on transcytosis and BBB permeability, will shed new light on the development of new therapeutic strategies. It has been postulated that Mfsd2a may serve to modulate transcytotic mechanisms in CNS endothelial cells for therapeutic purposes (Betsholtz, 2014; Zhao and Zlokovic, 2014). By pinpointing the specific transcytotic route that Mfsd2a suppresses, and by illuminating a novel role of membrane lipid metabolism in BBB function, this work may prompt the development of new strategies to deliver drugs to the diseased CNS, potentially via the use of lipid-based biologics to "de-repress" the caveolae pathway.

STAR*METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
 - Animal Models
 - Cell lines
- METHOD DETAILS
 - Lipid profiling
 - Immunohistochemistry and immunocytochemistry
 - BBB permeability assay
 - Transmission electron microscopy
 - Immunogold labeling for electron microscopy

- Isolation of capillary fragments
- Western blotting
- Lentivirus production
- In vitro cholera toxin subunit-B uptake assays
 Imaging
- QUANTIFICATION AND STATISTICAL ANALYSIS
 - Quantification of BBB leakage
 - TEM Quantification
 - Statistical Analysis
- DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures, four tables, and one data file and can be found with this article online at http://dx.doi.org/10.1016/j.neuron. 2017.03.043.

AUTHOR CONTRIBUTIONS

B.J.A. and C.G. conceived the project and designed experiments. K.B., A.A.D., and C.B.C. performed lipidomic mass spectrometry. B.J.A., A.T., and B.L. performed all other experiments. A.B.-Z. initiated lipidomic experiments. D.D.G. provided experimental resources. B.J.A. and B.W.C. analyzed all data. B.J.A. and C.G. wrote the manuscript, with input from all authors.

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KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit polyclonal α-Mfsd2a	Cell Signaling Technologies	Under Development; Cat #Mfsd2a14-87; RRID: AB_2617168
Mouse monoclonal α-Claudin-5-488 conjugate	Thermo Fisher Scientific	Cat #352588; RRID: AB_2532189
Rat polyclonal α-PECAM	BD Biosciences	Cat #553370; RRID: AB_394816
Rabbit polyclonal α-Cav-1	Cell Signaling Technologies	Cat #3267; RRID: AB_2275453
Rabbit polyclonal α-NeuN	Millipore	Cat #MAB377; RRID: AB_2298772
Rabbit polyclonal α-VE-cadherin	Abcam	Cat #ab33168; RRID: AB_870662
Rabbit polyclonal α -PDGFR- β	Cell Signaling Technologies	Cat #3169; RRID: AB_2162497
Rabbit polyclonal α-Aquaporin-4	Millipore	Cat #ABN411; RRID: AB_2637077
Mouse monoclonal α- β-Actin	Sigma	Cat #A2228; RRID: AB_476697
Mouse monoclonal α-Tubulin	Sigma	Cat# T5168; RRID: AB_477579
Chemicals, Peptides, and Recombinant Proteins		
Isolectin GS-IB4, from Griffonia simplicifolia	Thermo Fisher Scientific	Cat #I21411
10-kDa Dextran-tetramethylrhodamine conjugate	Thermo Fisher Scientific	Cat #D1817
Horseradish peroxidase, type II	Sigma-Aldrich	Cat #P8250-50KU
Cholera toxin subunit B, Alexa Fluor 555 conjugate	Thermo Fisher Scientific	Cat #C34776
PNA Bio Cas9 protein	Thermo Fisher Scientific	Cat #NC0789474
Experimental Models: Cell Lines		
Human: HEK293T cells	ATCC	CRL-3216
Mouse: bEnd.3 cells	ATCC	CRL-2299
Human: PANC-1-MSFD2A-GFP	Dr. David Sabatini; (Reiling et al., 2011)	N/A
Experimental Models: Organisms/Strains		
Mouse: Mfsd2a KO: B6;129S5-Mfsd2a ^{tm1Lex} /Mmucd	Mouse Biology Program, University of California, Davis	MMRRC strain 032467-UCD; RRID: MGI:5755244
Mouse: Cav-1 KO: STOCK Cav1 ^{tm1Mls} /J	Jackson Laboratory	JAX: 004585; RRID: IMSR_JAX:004585
Mouse: Mfsd2a ^{D96A}	This paper	N/A
Oligonucleotides		
Mfsd2a ^{D96A} gRNA: GCCTTCACTGACCCTCTGGTGGG	Integrated DNA Technologies	N/A
Mfsd2a ^{D96A} Donor Oligonucleotide: TGCTTCTTCCCCA TGGCAGGTGGAACCACTTCCTGCTTCCATTATCCTTT TTGTGGGCCGAGCCTGGGATGCCTTCACTGCCCCA CTAGTGGGCTTCTGCATTAGCAAGTCCTCCTGGACC CGCCTGGGCCGCCTCATGCCCTGGTAAGTAGAGAC GCTCCTTCGAAG	Integrated DNA Technologies	N/A
Genotyping Primers: Mfsd2a ^{D96A} : (Fwd): TTGGTGGTTC AGCTGTGCCA; (Rev): GACACCAGAAGCCTGCCTTT; (Sequencing): ATGCTCCGCTAAGGCAATGA	Integrated DNA Technologies	N/A
Genotyping Primers: Mfsd2a: (WT Fwd): CCTGGTTTG CTAAGTGCTAGC; (WT Rev): GTTCACTGGCTTGGAG GATGC; (KO Fwd): CACTTCCTAAAGCCTTACTTC; (KO Rev): GCAGCGCATCGCCTTCTATC	Integrated DNA Technologies	N/A

(Continued on next page)

Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Genotyping Primers: Cav1: (WT Fwd): GCACACCAA GGAGATTGACC; (KO Fwd): CTCCAGACTGCCTTG GGAAAA; (Rev): CTTGGCTGTCACCACACAC	Integrated DNA Technologies	N/A
Recombinant DNA		
pCAG-GFP	Matsuda and Cepko, 2004	Addgene Plasmid #11150
pHAGE-PGK_IZsGreen	Harvard Medical School, Jeng-Shin Lee	https://plasmid.med.harvard.edu/ PLASMID/GetVectorDetail. do?vectorid=251
Plasmid: pCAG- <i>mMfsd2a</i> -GFP	This paper	N/A
Plasmid: pHAGE-PGK_mMfsd2a-IZsGreen	This paper	N/A
Software and Algorithms		
Prism 7	GraphPad	http://www.graphpad.com/scientific- software/prism/
CCTop – CRISPR/Cas9 target online predictor	Center for Organismal Studies Heidelberg; (Stemmer et al., 2015)	http://crispr.cos.uni-heidelberg.de/
TraceFinder 3.2	Thermo Fisher Scientific	http://tools.thermofisher.com/content/ sfs/manuals/Man-XCALI-97598- TraceFinder-32-User-Clinical- ManXCALI97598-A-EN.pdf
Custom ImageJ macro for quantifying BBB permeability	This paper	Data S1

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Chenghua Gu (chenghua_gu@hms.harvard.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animal Models

All mice were group housed in standard vivarium conditions, with ad libitum access to diet and water. *Mfsd2a* null mice (MMRRC strain 032467-UCD; RRID: MGI:5755244) were maintained on a mixed B6/129 background and were used to test the involvement of Mfsd2a in transcytosis and BBB permeability. *Cav-1* null mice (JAX: 004585; RRID: IMSR_JAX:004585, mixed B6/129 background) were used to test the involvement of caveolae-mediated transcytosis in BBB permeability.

Mfsd2a^{D96A} mice were generated at HHMI Janelia Farm. The D96A mutation was generated using the following: gRNA sequence 5'-GCCTTCACTGACCCTCTGGTGGG-3', donor oligonucleotide sequence 5'-TGCTTCTTCCCCATGGCAGGTGGAAC CACTTCCTGCTTCACTGACCCTTCAGTGGGCCCGAGCCTGGGACCCGGGAGCCTGGGATGCCTTCACTGCCCCACTAGTGGGGCTTCTGCATTAGCAAGTCC TCCTGGACCCGCCTGGGCCGCCTCATGCCCTGGTAAGTAGAGAGACGCTCCTTCGAAG-3', and cas9 protein (Thermo Fisher Scientific, Cat #NC0789474), with an injection concentration: Cas9/gRNA/donor oligonucleotide = 100ng/100ng/20ng/µl. The mouse strain B6SJF1/J (JAX: 100012) was used for injection, with an injection efficiency of 62.5% (20/32). Founder *Mfsd2a^{D96A}* mice were backcrossed to C57BL/6 mice (Charles River) for three generations prior to analysis. The open-access online tool, CCTop (Stemmer et al., 2015) was used to analyze the possibility of off-target DSBs in *Mfsd2a^{D96A}* mice. The possibility of off-target mutations found in *Mfsd2a^{D96A}* genomic DNA. A 250 bp PCR product containing *Mfsd2a* exon 3 was amplified using the following primers: 5'- TTGGTGGTTCAGCTGTGCCA-3' and 5'- GACACCAGAAGCCTGCCTTT-3', and Sanger sequenced with 5'- ATGCT CCGCTAAGGCAATGA-3' to confirm generation of the D96A point mutation and Spel restriction site. For genotyping, PCR products were digested with Spel and run on agarose gels to test for the presence of the *Mfsd2a^{D96A}* allele.

No animal randomization was performed and both male and female mice were used for experiments and no sex-based differences in phenotype were observed. Animals were genotyped via allele-specific PCR reactions and experiments were performed with observer not blinded to genotype. Information regarding the age of mice used for each specific experiment can be found in the Results and Figure Legends. All animals were treated according to institutional and US National Institutes of Health (NIH) guidelines approved by the International Animal Care and Use Committee (IACUC) at Harvard Medical School.

Cell lines

HEK293T (ATCC, #CRL-3216) and bEnd.3 (ATCC, #CRL-2299) cells were cultured in DMEM supplemented with 10% FBS and 1% penicillin/streptomycin in 5% CO₂ at 37°C. Cell lines were obtained directly from ATCC, with no additional cell authentication performed. PANC-1 cells stably transfected with human MFSD2A-GFP (Reiling et al., 2011) were obtained from Dr. David Sabatini and cultured in DMEM supplemented with 10% FBS, 1% penicillin/streptomycin, and 2 μ g/ml puromycin in 5% CO₂ at 37°C.

METHOD DETAILS

Lipid profiling

Lipids from whole brain, and brain and lung isolated capillary lysates were profiled using liquid chromatography tandem mass spectrometry (LC-MS) method operated on a Nexera X2 U-HPLC (Shimadzu Scientific Instruments) coupled to an Exactive Plus orbitrap MS (Thermo Fisher Scientific). Briefly, tissue samples were homogenized in a weight proportional volume of water (4µl/mg of tissue) using a TissueLyser II (QIAGEN). Equal starting amounts of tissue homogenate were used as starting material for all mass spectrometry. Lipids were extracted from homogenates (10 µl) using 190 µl of isopropanol containing 1,2-didodecanoyl-sn-glycero-3-phosphocholine as an internal standard (Avanti Polar Lipids). After centrifugation (10 min, 10,000 x g, ambient temperature), supernatants (2 µl) were injected directly onto a 100 × 2.1 mm ACQUITY BEH C8 column (1.7 µm; Waters). The column was eluted at a flow rate of 450 μL/min isocratically for 1 min at 80% mobile phase A (95:5:0.1 vol/vol/vol 10 mM ammonium acetate/methanol/formic acid), followed by a linear gradient to 80% mobile-phase B (99.9:0.1 vol/vol methanol/formic acid) over 2 min, a linear gradient to 100% mobile phase B over 7 min, and then 3 min at 100% mobile-phase B. MS analyses were carried out using electrospray ionization in the positive ion mode using full scan analysis over m/z 200-1100 at 70,000 resolution and 3 Hz data acquisition rate. Additional MS settings were: ion spray voltage, 3.0 kV; capillary temperature, 300°C; probe heater temperature, 300°C; sheath gas, 50; sweep gas, 5; and S-lens RF level 60. Raw data were processed using TraceFinder 3.2 software (Thermo Fisher Scientific) to integrate extracted ion chromatograms for approximately 240 lipids characterized by headgroup and total acyl chain carbon and double bound content. For data analysis, all individual lipid levels (integrated peak area) were normalized to internal standard values prior to statistical testing. Average values of integrated peak area were used for comparisons between conditions for each lipid species.

Immunohistochemistry and immunocytochemistry

Mouse tissues were fixed by immersion in 4% PFA/PBS overnight at 4°C, cryopreserved in 30% sucrose, and frozen in TissueTek OCT (Sakura). Tissue sections were blocked with 10% goat serum/5% BSA/PBST (0.5% Triton X-100) and stained overnight at 4°C with the following primary antibodies: α -Mfsd2a (1:200, Cell Signaling Technologies; RRID: AB_2617168), α -PECAM (1:200, BD Biosciences #553370; RRID: AB_394816), α -Claudin-5-488 conjugate (1:300, Thermo Fisher Scientific #352588; RRID: AB_2532189), isolectin B4 Alexa 488 conjugate (1:200, Thermo Fisher Scientific #121411), followed by corresponding Alexa Fluor-conjugated secondary antibodies (1:1000, Thermo Fisher Scientific). Sections were mounted with ProLong Gold for imaging. bEnd.3 cells were fixed in 4% PFA/PBS for 15 min at room temperature as processed as described above.

BBB permeability assay

Postnatal

P4-P5 pups were anaesthetized with isoflurane and 10-kDa dextran tetramethylrhodamine (Thermo Fisher Scientific, Cat #D1817, 10 mg/ml, 10 μ L per g body weight) was injected into the left heart ventricle with a Hamilton syringe. After five minutes of circulation, brains were dissected and fixed by immersion in 4% PFA/PBS at 4°C overnight, cryopreserved in 30% sucrose, and frozen in TissueTek OCT (Sakura). Sections of 14 μ m were collected and co-stained with α -Claudin-5-488 conjugate (1:300, Thermo Fisher Scientific #352588; RRID: AB_2532189) or isolectin B4 Alexa 488 conjugate (1:200, Thermo Fisher Scientific #121411) to visualize blood vessels.

Adult

Adult P90 mice were briefly anesthetized with isoflurane and HRP type II (Sigma Aldrich, Cat #P8250-50KU, 0.5 mg/g body weight dissolved in 0.4 mL PBS) was injected bi-laterally into the retro-orbital sinus. After 30 min of circulation, brains were dissected and fixed by immersion in 4%PFA/PBS at 4°C overnight. Following fixation, brains were washed in PBS and forebrain vibratome free-floating sections of 50 µm were collected, processed in a standard DAB reaction for 35 min, and mounted on 3% gelatin-coated slides. Slides were dehydrated through an ethanol series and mounted with Permount for brightfield imaging.

Transmission electron microscopy

In vivo

Brains from P4-P5 pups were dissected and fixed by immersion in 5% glutaraldehyde/4% PFA/0.1 M sodium-cacodylate for 10-14 days at room temperature. P90 mice were anaesthetized and HRP type II (Sigma Aldrich, Cat #P8250-50KU, 0.5 mg/g

body weight dissolved in 0.4 mL PBS) was injected bilaterally into the retro-orbital sinus. After 30 min of circulation, brains were dissected and fixed by immersion, first in 5% glutaraldehyde/4% PFA/0.1 M sodium-cacodylate for 1 hr at room temperature, then overnight at 4°C in 4% PFA/0.1 M sodium-cacodylate. Following fixation, brains were washed overnight in 0.1 M sodium-cacodylate. Coronal vibratome free-floating sections of 50 μ m were collected, post-fixed in 1% osmium tetroxide and 1.5% potassium ferrocyanide, dehydrated, and embedded in epoxy resin. Ultrathin sections of 80 nm were then cut from the block surface, collected on copper grids, and counter-stained with Reynold's lead citrate.

In vitro

Experiments on bEnd.3 cells were performed 5 days post-infection with lentivirus. bEnd.3 and PANC-1 confluent cell cultures on aclar coverslips were fixed in a 2.5% paraformaldehyde/5% glutaraldehyde/0.06% picric acid/0.2 M sodium-cacodylate mixer diluted 1:1 in culture media for 15 min at room temperature. Cultures were processed for TEM, as described above.

Immunogold labeling for electron microscopy

Adult P90 mice were anesthetized and perfused through the heart with 30 mL PBS, followed by first 150 mL of a 5% glutaraldehyde/ 4% PFA/0.1 mM phosphate buffer fixative solution, then a 4% PFA/0.1 mM phosphate buffer fixative solution. Brains were dissected and post-fixed in 4% PFA/PBS for 30 min at 4°C. Coronal vibratome free-floating sections of 50 μ m were collected and immersed in 0.1% sodium borohydride/PBS for 20 min at room temperature, blocked with 10% goat serum/0.5% gelatin/PBST (0.01% Triton X-100), and incubated with α -Cav-1 (1:100, Cell Signaling Technologies #3267; RRID: AB_2275453) overnight at room temperature. Sections were then incubated with gold-labeled goat α -rabbit IgG (1:50, Nanoprobes) overnight at room temperature, washed with PBS and sodium acetate, and silver-enhanced prior to processing for TEM.

Isolation of capillary fragments

Capillary isolation procedure was adapted from a combination of previously described protocols (Goldstein et al., 1984). P4 mouse brains were dissected and homogenized in PBS. Homogenates were centrifuged for 30 min on a density gradient of 16% dextran (Sigma-Aldrich). The tissue pellet containing capillary fragments was resuspended in PBS and filtered through a 35 µm cell strainer to remove red blood cells and cell-type contaminates. P4 mouse lungs were processed as described above, with an additional tissue dissociation step using Collagenase/Dispase (Roche) for 30 min at 37°C prior to centrifugation on a dextran gradient. Isolated capillary fragments were either snap frozen in liquid nitrogen for lipid extraction or lysed and processed for western blotting.

Western blotting

Protein lysate samples were boiled at 95°C, run on 6%–12% polyacrylamide gels, and electrophoretically transferred onto PVDF membranes. Membranes were blocked with 5% BSA/TBST (0.1% Tween-20) or 5% non-fat milk/TBST and incubated overnight at 4°C with the following primary antibodies: α -Mfsd2a (1:1000, Cell Signaling Technologies; RRID: AB_2617168), α -NeuN (1:2000, Millipore #MAB377; RRID: AB_2298772), α -VE-Cadherin (1:1000, Abcam #ab33168; RRID: AB_870662), α -Aquaporin-4 (1:1000, Millipore #ABN411; RRID: AB_2637077), α -PDGFR- β (1:1000, Cell Signaling Technologies #3169; RRID: AB_2162497), α - β -Actin (1:5000, Sigma Aldrich #A2228; RRID: AB_476697), and α -Tubulin (1:8000, Sigma Aldrich #T5168; RRID: AB_477579). Membranes were then incubated with the appropriate HRP-conjugated secondary antibody (1:10000, Southern Biotech) and developed with regular or super ECL (Thermo Fisher Scientific). The intensity of individual bands was quantified using ImageJ.

Lentivirus production

HEK293T cells were transfected using Lipofectamine 2000 (Thermo Fisher Scientific) with the pHAGE-PGK_IZsGreen lentiviral plasmid containing full-length mouse *Mfsd2a* CDS, along with Δ-vpr and vsv-g packaging plasmids. Supernatant was harvested after three days and concentrated overnight at 4°C with Lenti-X Concentrator (Clontech). Lentivirus was stored fresh at 4°C for up to 24 hr prior to use.

In vitro cholera toxin subunit-B uptake assays

Uptake procedure was adapted from a combination of previously described protocols (Kirkham et al., 2005). Confluent bEnd.3 cells were transfected using Lipofectamine 2000 (Thermo Fisher Scientific) with either pCAG-*GFP* or pCAG-*mMfsd2a*-*GFP* for two days and serum-starved for two hours prior to experiments. Cultures were then incubated with cholera toxin subunit-B (CTB) Alexa-555 conjugate (Thermo Fisher Scientific, Cat #C34776, 150 µg/ml) for ten minutes at 37°C, washed several times with cold PBS, fixed in 4% PFA/PBS for 15 min at room temperature, washed with PBS, and mounted with ProLong Gold for imaging.

Imaging

An Olympus FluoView FV1000 or FV1200 laser scanning confocal microscope was used to image dextran tracer permeability assays and expression analyses. A Nikon Ti spinning disk confocal microscope was used to image the in vitro CTB uptake assay. An Olympus VS120-SL 5 slide scanner was used to image adult HRP tracer permeability assays. A 1200EX electron microscope (JOEL) equipped with a 2k CCD digital camera (AMT) was used for all TEM studies. Images were processed using ImageJ (NIH) and Adobe Photoshop.

QUANTIFICATION AND STATISTICAL ANALYSIS

Quantification of BBB leakage

Postnatal dextran tracer permeability assay

All quantification was performed blind using a custom ImageJ (NIH) macro (see Data S1). 6-8 images per animal were obtained from 14 µm thick coronal forebrain sections of the same rostrocaudal position using a 40x oil immersion objective. The same forebrain regions were imaged among littermate pairs. The same background subtraction and auto threshold parameters were applied to max z-projections of both Isolectin/Claudin-5 (endothelium vessel markers), and dextran (tracer) channels. Masks of both channels were generated and the permeability index of leakage was defined as the area of tracer mask divided by area of vessels mask. A leakage value of 1 signifies no tracer extravasation, while significantly higher values signify tracer extravasation.

TEM Quantification

For all TEM quantifications, mean vesicular pit/vesicular density values were calculated from the number of vesicles per μ m of membrane for each image collected. All images were collected at 12000x magnification. Density values for each cell source/genotype reflect the average of mean density values obtained from each individual experiment/animal and are expressed as mean ± s.e.m.

Statistical Analysis

All statistical analyses were performed using Prism 7 (GraphPad Software). Two group comparisons were analyzed using an unpaired two-tailed Student's t test. Multiple group comparisons were analyzed using a one-way ANOVA, followed by a post hoc Tukey's test. Sample size for all experiments was determined empirically using standards generally employed by the field, and no data was excluded when performing statistical analysis. Standard error of the mean was calculated for all experiments and displayed as errors bars in graphs. Statistical details for specific experiments, including exact n values and what n represents, precision measures, statistical tests used, and definitions of significance can be found in the Figure Legends.

DATA AND SOFTWARE AVAILABILITY

A custom ImageJ macro was used to quantify postnatal dextran tracer permeability assays, as described in the Method Details. For script, see Data S1.