Over-expression of Slit2 induces vessel formation and changes blood vessel permeability in mouse brain

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Aim: To investigate the effect of the axon guidance cue Slit2 on the density of blood vessels and permeability of the blood-brain barrier in mouse brain.

Methods: hSlit2 transgenic mouse line was constructed, and the phenotypes of the mice were compared with wild-type mice in respect to the lateral ventricle (LV), ventricle pressure, and the choroids plexus. An in vivo Miles permeability assay and an amyloid-β permeability assay were used to assess the permeability of brain blood vessels. Brain vessel casting and intracerebral hemorrhage models were built to investigate vessel density in the transgenic mice. An in vitro permeability assay was used to test whether Slit2 could change the permeability and tight junctions of blood vessel endothelial cells.

Results: Hydrocephalus occurred in some transgenic mice, and a significantly larger lateral ventricle area and significantly higher ventricle pressure were observed in the transgenic mice. The transgenic mice displayed changed construction of the choroids plexus, which had more micro vessels, dilated vessels, gaps between epithelial cells and endothelial cells than wild-type mice. Slit2 significantly increased brain vessel density and the permeability of brain vessels to large molecules. These blood vessels were more sensitive to cues that induce brain hemorrhage. At the cellular level, Slit2 disturbed the integrity of tight junctions in blood vessel endothelial cells and improved the permeability of the endothelial cell layer. Thus, it promoted the entry of amyloid-β peptides from the serum into the central nervous system, where they bound to neurons.

Conclusion: Slit2 increases vessel density and permeability in the brains of transgenic mice. Thus, Slit2 induces numerous changes in brain vessels and the barrier system.

Keywords: Slit2; blood-brain barrier; permeability; vessel endothelial cell; tight junction

Introduction

Vasogenic brain edema, which is defined as the translocation of proteins and fluid from the vascular space across the blood-brain barrier (BBB)[1], is a major life-threatening complication of various injuries to the central nervous system (CNS)[2,3]. Endothelial cells of the brain vasculature form the BBB and maintain the homeostasis of the central nervous system (CNS). Pathological conditions such as brain tumors and head injuries increase the permeability of the brain microvasculature and destroy the BBB[4]. Vascular endothelial growth factor (VEGF) is well known as the major inducer of angiogenesis, and it also increases the permeability of the microvasculature and stimulates endothelial cell growth[5-8]. Increased VEGF expression may cause vascular leakage in the CNS in vivo[9]. However, the underlying molecular and pathogenic mechanisms behind edema and blood-brain vessel leakage are poorly understood.

The Slit family of guidance cues interacts with the Roundabout (Robo) family of transmembrane receptors in physiological and pathological processes requiring cell migration[10-13]. During development of the nervous system, Slit-Robo signaling regulates the repulsion or attraction of projecting axons and migrating neurons[10-13]. Vascular endothelial cells secrete Slit2, which binds to Robo1 on leukocytes and acts as an endogenous inhibitor of leukocyte chemotaxis[16-21]. Additionally, Slit2 mediates directional migration of malignant cells[22-24]. We and others have previously reported that Slit proteins secreted by solid tumors bind to Robo1, which is expressed on vascular and lymphatic endothelial cells, to
stimulate angiogenesis and lymph angiogenesis\textsuperscript{25–31}. Slit2 is expressed in the CNS while Robo1 is expressed in blood vessel endothelial cells. However, whether expression of these proteins can change the permeability of blood vessels and whether abnormal expression can induce vessel leakage and edema remain to be determined.

In this study, we constructed a transgenic mouse line that over-expresses human Slit2 and observed that hydrocephalous occurs in some of these transgenic mice. The transgenic mice also had larger lateral ventricles and higher ventricle pressure than wild-type mice. Comparison of the choroid plexus, where cerebrospinal fluid (CSF) is secreted, revealed that there was a change in the construction of the choroid plexus, with the transgenic mice having more microvessels, dilated vessels, gaps between epithelial cells and endothelial cells. We also found that Slit2 could improve brain vessel density and promote the permeability of brain vessels to large molecules. These blood vessels were also more sensitive to cues that induced brain hemorrhage. At the cellular level, Slit2 disturbed the integrity of tight junctions in blood vessel endothelial cells and increased the permeability of the endothelial cell layer. The ability of Slit2 to increase the permeability of the BBB resulted in an increase in the transfer of amyloid-β peptides from the serum to the CNS, where they bound to neurons.

Material and methods

Generation of hSlit2 transgenic mice and detection of hSlit2 over-expression

hSlit2 transgenic mice were generated according to standard procedures. The transgene was constructed by cloning cDNA encoding full-length human Slit2 between the BamH I and Xho I restriction sites of the MCS (multi clone site) of pCEP4F. Genotypes were confirmed by Southern blot and PCR analysis. PCR screening of hSlit2 heterozygotes was performed on standard tail genomic DNA preparations using a pair of primers specific for human Slit2 cDNA (forward: 5'-GGTGACCGGATC-CCATATCGCGGTAGAACTC-3'; reverse: 5'-GGACACCTC-GAGCGTACAGCCGCACTTCAC-3'). PCR cycles were as follows: 95°C, 4 min (1 cycle); 94°C, 45 s; 55°C, 45 s; and 72°C, 1 min (63 cycles); and 72 °C, 10 min (2 cycles). PCR products were analyzed on 1% agarose gels. Slit2 homozygosity was confirmed by genetic methods based on the principle that the progeny of Slit2 homozygotes mated to wild-type C57 mice should all be heterozygotes. The brains from C57 control littermate mice and hSlit2 transgenic mice from founder #9 were snap frozen in liquid N\textsubscript{2} and pulverized. The brain powder was homogenized in 1 mL RIPA lysis buffer [50 mmol/L Tris-HCl, pH 7.4, 150 mmol/L NaCl, 1% Nonidet P-40, 0.5% deoxycholic acid, 0.1% sodium dodecyl sulfate (SDS), 5 mmol/L EDTA, 2 mmol/L phenylmethylsulfonyl fluoride (PMSF), 20 μg/mL aprotinin, 20 μg/mL leupeptin, 10 μg/mL pepstatin A, and 150 mmol/L benzamidine] in a Dounce tissue homogenizer. After homogenization, the samples were centrifuged at 12000 × g for 10 min to remove tissue debris and boiled in SDS sample buffer for 5 min. They were then subjected to 7% SDS-PAGE electrophoresis, transferred to blotting membranes, probed with 1 μg/mL anti-Slit2 monoclonal antibody (5A5) and detected with the horseradish peroxidase-conjugated goat anti-mIgG Ab using a chemiluminescent detection system.

Comparison of the LV area and the choroid plexus

The mouse brains were fixed in 4% PFA (Sigma) and cut into consecutive longitudinal sections. A photo was taken of every longitudinal sections from each brain with an Olympus MVX10. Then the area of the lateral ventricle was calculated by Image Tool 3 (UTHCSA), and the average area of consecutive sections from each mouse was calculated. The data represent the means for groups of six mice. For histological examination of the choroid plexus, brains were isolated from age-matched C57 and Slit2 adult mice, cut into small coronal blocks, fixed in 4% formaldehyde, embedded in paraffin, and cut into 0.5-mm sections following standard procedures. Tissue sections were counterstained with hematoxylin. The same regions of the choroid plexus in all mouse brains were photographed.

Electron microscopy

The choroid plexuses used for TEM were fixed in 2% glutaraldehyde and 1% sucrose in 0.1 mol/L cacodylate buffer for 3 h. The samples were washed in 0.1 mol/L cacodylate buffer and then post-fixed in 1% osmium tetroxide in 0.1 mol/L cacodylate buffer for 2 h, all at pH 7.0 and 25°C. The choroid plexuses were embedded in epon after dehydration in a graded series of ethanol. Epon was polymerized at 60°C for 48 h. Serial sections of 80 nm were cut on a Leica Super Nova ultramicrotome with a diamond knife and collected on formvar-coated nickel grids. Sections were contrasted with uranylacetate and stained with 1% toluidine blue. The sample grids were observed using a Hitachi 600 TEM.

In vivo Miles permeability assay and amyloid-β permeability assay

To investigate the effects of over-expression of hSlit2 in the brain on vascular permeability, a Miles assay was performed. Mice received an iv injection of sterile 0.5% Evans blue dye (200 μL) via the tail vein. Mice were killed 20 min after the injections by cervical dislocation after anesthesia. The brains (weighted) were cut into small pieces and incubated in 500 mL of formamide at 37°C for 48 h to extract the Evans blue dye. The absorbance of the extracts was read at 630 nm in a spectrophotometer (Beckman DU 640). For the amyloid-β permeability assay, mice received an iv injection of 6.9 mmol/L FITC-Aβ (100 μL) via the tail vein. Mice were killed 48 h after the injections by cervical dislocation after anesthesia. The brains of the mice were prepared on crystal slides, and the nuclei were stained with DAPI.

LV pressure assay

Mice (12 weeks of age) were anesthetized by ip injection of sodium pentobarbital (70 mg/kg). A small hole 0.5 mm posterior and 1.0 mm lateral of the bregma was drilled to perfo-
rate the skull. A pressure transducer linked to a monitor was injected into the brain at a depth of 2.3 mm unilaterally. Then the LV pressure was read on a computer that was linked to the monitor (Powerlab 4/30). The pressure was measured as mmH2O.

Immunohistochemical staining
Antibodies against vWF (Antibody Diagnostica Inc; a 1:200 dilution for paraffin-embedding), Slit2 (5 μg/mL for paraffin-embedding sections), and Robo1 (20 μg/mL for paraffin-embedding sections) were used for immunohistochemical staining as described previously.

Brain vessel casting
After systemic heparinization with 750 IU/kg intravenous heparin, the common carotid arteries were cannulated and perfused with approximately 100 mL of 27 °C saline, followed by a 2.5% buffered glutaraldehyde solution (Sigma) at pH 7.4. The casts were made by perfusion of the arteries with 100 mL of Mercox (SPI, West Chester, PA, USA) diluted with 20% methyl methacrylate monomers (Aldrich Chemical, Milwaukee, WI, USA). After complete polymerization, the brains were harvested and macerated in 5% potassium hydroxide, followed by drying and mounting for scanning electron microscopy. The microvascular corrosion casts were imaged after being coated with gold with a Hitachi S-450 scanning electron microscope.

Intracerebral hemorrhage (ICH) model
Mice (12 weeks of age) were prepared for surgery and anesthetized by ip injection of sodium pentobarbital (70 mg/kg) [32]. A small hole 1.0 mm posterior and 3.0 mm lateral of the bregma was drilled to perforate the skull. A 1-μL Hamilton syringe was used to deliver 500 nL of collagenase/saline (150 U/μL) to the caudate/putamen at a depth of 4.0 mm unilaterally. After the injection of collagenase/saline (~30 s), the needle remained in place for another 2 min to prevent reflux of fluid. Then the scalp skin was closed using 4.0 nylon sutures. Twenty-four hours later, the mice were perfused with PBS, the brains were harvested, and 14-μm sections were prepared using a cryostat and mounted on glass slides. An Olympus BX100 upright systems microscope with a digital camera was used to capture images. The hemorrhage volume was measured using the Stereologer software system.

In vitro permeability assay
We coated transwell inserts (Corning, 48-well, 3-mm pore) with collagen and seeded HUVEC cells at a density of 30000 cells per well. Cells were then cultured for another 24 h. One hour later, Slit2, VEGF, R5, and FITC-DEXTRAN (25 mg/mL, Sigma) were added to the top of the inserts. The absorbance of the solution in each well was measured at 492 nm (n=6 wells each).

Statistical analysis
Statistical significance was determined by Student’s t-test. P-values of 0.05 and 0.01 were considered statistically significant and very significant, respectively.

Results
Generation of hSlit2 transgenic mice and over-expression of hSlit2 in mouse brains
To study the function of Slit2 in the whole blood vessel system, we constructed a Slit2 over-expressing mouse transgenic plasmid, for which a schematic display is shown in Figure 1A. The full-length human Slit2 cDNA, which is 4689 bases, was inserted between the BamHI/XhoI restriction sites of the MCS of the pCEP4F vector, which has a pCMV promoter and an N-terminal flag-tag. The plasmid was linearized with restriction enzymes, injected into the pronuclei of fertilized C57×CBA F1 oocytes and transplanted into the mother mouse. The offspring mice were analyzed by dot blot (Figure 1B). The presence of the transgene in mice was verified by dot blotting using a [32P]-labeled fragment of the hSlit2 plasmid. (C) Southern blotting of hSlit2 transgenic mice. The presence of the transgene was further confirmed by Southern blotting of DNA isolated from mice that were shown to be positive for the transgene based on the dot blot analysis with a [32P]-labeled fragment of the hSlit2 plasmid. (D) Expression of the hSlit2 protein in the hSlit2 transgenic mice compared with C57 mice. The expression of Slit2 (~200 kDa) in the brains of C57 and Slit2 mice was detected with anti-Slit2 IgG 5A5. (E) RT-PCR of hSlit2 in mRNA extracts from C57 and hSlit2 mouse brains. The expression of Slit2 (~200 kDa) in the brains of C57 and Slit2 mice was detected with anti-Slit2 IgG 5A5. (F) Western blot analysis of flag-tagged proteins. Brain lysates from C57 and hSlit2 mice that were ip injected with an anti-Slit2 antibody, 10D10, and probed with an anti-flag tag antibody.
Mouse lines 9, 25, and 47 were shown to express human Slit2. The presence of the human Slit2 transgene was also confirmed by Southern blot analysis (Figure 1C). Among the three transgenic strains, we found that strain 9 had the strongest signal in dot blot and Southern blot assays, so we focused our research on strain 9. We observed that the transgenic mice expressed more Slit2 in the brain compared with C57 mice at the transcriptional and translational levels (Figure 1D, 1E). We used the Slit2 antibody 10D10 to immune-precipitate the mouse brain lysate and detected the precipitate with anti-flag antibody, and we found that flag-Slit2 was expressed in the transgenic mouse brain (Figure 1F).

**The phenotype of Slit2 mice and comparison of the LV area**

Most of the transgenic mice initially appeared normal (Figure 2A), but about 5% of the transgenic mice had an intumescent head (Figure 2B). This phenotype appeared at the age of 4–6 weeks in all three of the transgenic mouse lines, line 9, line 25, and line 47. Mice with an intumescent head died within 2 weeks, at the age of 6–8 weeks. Anatomical analysis of these mice revealed high levels of encephalopathic edema. In mice where this brain phenotype was observed, the brain edema was serious and the brain tissue was destroyed to such a degree that the structure of the brain could not be recognized. Thus, we focused our research on the transgenic mice that did not have this phenotype. Brain edema is linked to the CSF system, including the production, circulation and absorption of CSF. We therefore examined brain structures related to the CSF system. In the brain, CSF is produced in the lateral ventricle, so we compared the lateral ventricle of transgenic mice and non-transgenic mice. We found that the lateral ventricle area was larger in the brains of transgenic mice (Figure 2C, 2D) than in control mice (Figure 2E, 2F). For this study, the mice were all 8 weeks old, and the same result was seen in male and female mice. Figure 2G shows a schematic display of the mouse brain. The arrow indicates the lateral ventricle. We calculated the entire area of the LV in slides created from the brains of each mouse with Image Tool 3 (UTHSCSA) and found that the LV area was larger in transgenic mice (n=6, P<0.01) (Figure 2H). LV pressure was also detected in the lateral ventricles of the mouse brains. In transgenic mice, the pressure was 128 mmH2O, and in control mice, it was 81 mmH2O (Figure 2I) (n=7, P<0.01). Thus, the LV pressure was higher in the brains of transgenic mice compared with control mice.

**Changes in the structure and function of the choroid plexus**

In mouse brains, the choroid plexus produces CSF. The structure of the choroid plexus was obviously different between transgenic and control mice. In control mice, it was a tight, monolayer covered with microvessels (Figure 3A); however, in transgenic mice, there were gaps between the epithelial cell layer and the vessels (Figure 3B, 3C), the vessels were enlarged and had gaps, and the epithelial cells were crenated (Figure 3C). The mice used in these experiments were all 10 weeks old. Examination under an electron microscope revealed gaps between the epithelial tight junctions in transgenic mice (Figure 3E); such gaps were not observed in control mice (Figure 3D). ZO-1 is an important component of cell-cell tight junctions and can be used as a marker for the completeness and density of tight junctions. We used fluorescence
immunostaining to detect ZO-1 in the choroid plexus. In control mice, ZO-1 had an equal and continuous distribution (Figure 3F), while in transgenic mice, the distribution of ZO-1 was disturbed (Figure 3G) and the tight junctions were destroyed.

More vessels are present in the choroid plexus and the brain of transgenic mice
Staining of slides of the brain cortex with anti-CD31 antibody showed that more micro-blood vessels were present in the brains of hSlit2 transgenic mice (Figure 4B) compared with C57 mice (Figure 4A). The gray density was 1.7% in C57 mice and 4.6% in hSlit2 transgenic mice (P<0.01, Figure 4C). We also stained the brain cortex slides with anti-vWF antibody and found that hSlit2 transgenic mice had an increased number of vessels (Figure 4E) compared with C57 mice (Figure 4D). Calculating the numbers of vessels present on the slides revealed that C57 mice had about 200 micro-vessels/mm², while hSlit2 transgenic mice had about 450 micro-vessels/mm² (Figure 4F), and most of these brain micro-vessels consisted of only one endothelial cell. Furthermore, we performed brain vessel casting, which also showed that hSlit2 transgenic mice had more brain vessels (Figure 4H) compared with C57 mice (Figure 4G).

Slit2 changes the permeability of cell-cell adhesions
To evaluate whether Slit2 improves the permeability of blood vessels, we performed a Miles assay on hSlit2 transgenic mice. An increased amount of Evans blue dye was detected in the brain tissue of transgenic mice compared with C57 mice (Figure 5A), which indicates that the blood vessels in the brains of transgenic mice are more permeable, allowing the entry of Evans blue dye into the tissue. Previous research has indicated that hyperplastic vessels are incomplete, more permeable, and more sensitive to destructive mechanism compared with normal vessels. Thus, we induced intracerebral hemorrhaging in the transgenic mice. This was accomplished by injecting collagenase into the mouse brains, which destroys blood vessels in the brain and causes intracerebral hemorrhaging. Examination of brain slides from mice in which intracerebral hemorrhaging was induced showed that transgenic mice have a larger hemorrhage area compared with the control mice (Figure 5C, 5D). Next, we measured the hemorrhage volume using the Stereologer software system and found that the hemorrhage volume also was increased in the transgenic mice (Figure 5B). These results indicate that the blood vessels in the brains of the transgenic mice are more permeable and much more sensitive to collagenase, which destroys the vessel structure. Thus, Slit2 may not only promote angiogenesis but could also increase the permeability of blood vessel endothelial cells, which are the most important component of blood vessels. VE-cadherin is a member of the cadherin family, which is expressed specifically in endothelial cells and plays important roles in endothelial cell cell-cell adhesion. Immunofluorescent detection of HUVECs using an anti-VE-Cadherin antibody revealed that, without incubation with Slit2, VE-Cadherin expression was equal, continuous and mostly localized to the conjunction sites of cell-cell adhesions (Figure 5E), while HUVECs incubated with the Slit2 protein had a disturbed distribution and lacked continuous expression of VE-Cadherin (Figure 5F). To verify that Slit2 can affect the permeability of blood vessels, we performed an in vitro permeability assay. We coated transwell inserts (Corning, 48-well, 3-mm pore) with collagen and seeded HUVECs at a density of 30000 cells per well. Once these cells formed a monolayer, we added Slit2 protein and other stimulating factors to the upper well. FITC-dextran was added 1 h later, and fluorescence was detected in the bottom wells. Our results show that VEGF-A can improve the permeability of HUVECs and Slit2 by 1.5 fold and 2 fold, respectively, and that a blocking antibody against the Slit2-Robo signal, R5, can block the effect of Slit2 on cell

Figure 3. Changes in the structure and the function of the choroid plexus. (A, B and C) The structure of the choroid plexus of a C57 mouse (A), a hSlit2 mouse (B) and a hSlit2 mouse with megahead (C). hSlit2 mice had structurally different choroid plexuses compared with C57 mice. Scale bar: 150 μm. (D and E) Electron micrograph of choroid plexus epithelial cells from the mouse lateral ventricle. Some vacuoles appear between the junctional complexes linking the epithelial cells in hSlit2 mice. Scale bar: 1.5 μm. (F and G) Immunofluorescence of the choroids plexus probed with an anti ZO-1 antibody. Cell-cell adhesion was different in hSlit2 mice compared with C57 mice.
permeability, returning it to basal levels (Figure 5G).

The presence of amyloid-β peptide in the brains of Slit2 mice
Amyloid-β 40 peptides were detected on slides of the transgenic mouse brains (Figure 6A) but not slides of C57 mouse brains (Figure 6D). In the cortex and hippocampus, amyloid-β 42 was detected in granal and pyramid cells in transgenic mice (Figure 6B, 6C) but not in C57 mice of the same age (Figure 6E, 6F). Furthermore, we injected FITC-Amyloid-β40 into the circulation by ip to see if the circulating peptides could enter the CNS. Florescence was observed on brain cortex slides prepared from the transgenic mice 1 h (Figure 6G), 24 h (Figure 6H), and 48 h (Figure 6I) after injection. The florescence was higher with increasing time, but even 48 h after injection, no florescence was detected on brain cortex slides prepared from C57 mice (Figure 6J, 6K).

hSlit2 transgenic mice have normal aqueduct and subarachnoid space, and Slit2 over-expression does not alter VEGF expression levels
To determine the course of the enlarged lateral ventricles observed in Slit2 transgenic mice, we compared the aqueduct and the subarachnoid space of transgenic and C57 mice. However, there were no obvious differences in these two areas between transgenic and C57 mice. The aqueduct was smooth and clear, and the aqueduct tube had the same inside diameter and did not contain any clogs (Figure 7A–7D). The SAS also had a complete and clear structure with no clogs or signs of collapse in the transgenic mice (Figure 7E–7H). We also found that Slit2 over-expression did not alter VEGF expression at either the protein or mRNA level (Figure 7I, 7J). These results show that the structures linked to circulation and absorption of CSF were complete and normal in the transgenic mice and that the enlargement of the lateral ventricle is caused by the abnormal production of CSF in the choroid plexus. On the other hand, the normal circulation and absorption of CSF could compensate for the abnormal production of cerebrospinal fluid, which could explain why edema was only observed in a small percentage of the transgenic mice.

Discussion
The brain barriers, including the blood-brain barrier, the blood-CSF barrier and the ventricular wall, provide a stable micro-environment for the proper functioning of the central nervous system. At the bases of the barrier structures are the junction structures, such as adherence junctions and tight junc-

Figure 4. More vessels are present in the choroid plexus and the brain of transgenic mice. (A, B and C) ICH of CD31 in C57 (A) and hSlit2 mice (B). Gray density statistics of CD31 staining (C), *P<0.01. Scale bar: 150 μm. (D, E and F) ICH of vWF in C57 (D) and hSlit2 mice (E). The difference in the number of stained vessels in the same area was statistically significant (F), *P<0.01. Scale bar: 50 μm. (G and H) Scanning electron microscopy of corrosion casts of the brain vessel of C57 (G) and hSlit2 mice (H).
tions between endothelial cells, epithelial cells and pericytes[33]. These junction structures are dynamic structures that consist of transmembrane proteins, cytoplasmic accessory proteins and scaffold proteins. Under different physiological and pathological conditions, changes occur in the expression, distribution, modification and interaction of these proteins[34]. These changes are regulated by several cell signaling pathways, and to date, the calcium channel pathway, the phosphorylation signaling pathway and the G-protein signaling pathway have been shown to change the expression and distribution of junction proteins, further affecting the function of barrier structures[35]. The brain blood vessel system is the structural basis of the blood-brain barrier, and the junctions between vessel endothelial cells have the most important effect on the permeability of the BBB. Many molecules, such as small chemical molecules, signaling proteins and inflammatory factors, can change the permeability of blood vessels. VEGF is an important angiogenic cue, and some reports have shown that VEGF improves the permeability of blood vessels and the BBB. VEGF binds to its receptor on blood vessel endothelial cells and triggers signaling pathways in the cell cytoplasm. This, in turn, alters the expression, phosphorylation and distribution of VE-cadherin and thus changes the permeability of blood vessels by disturbing the junction structures[36]. However, barrier structures are very complex and have many components. Therefore, whether other cues that induce angiogenesis have some effect on the permeability of the blood-brain barrier should be investigated.

The guidance cue Slit2 has been reported to regulate a number of physiological processes, mostly in the central nervous system, by controlling cell migration. Previous research from our lab indicated that Slit2 promotes tumor angiogenesis in a manner similar to VEGF[25]. In this paper, we found that Slit2 improved blood vessel density in the brain and promoted the permeability of brain blood vessels to large molecules. In addition, these blood vessels were more sensitive to cues that can induce brain hemorrhage. At the cellular level, Slit2 disturbed the integrity of blood vessel endothelial cell tight junctions and improved the permeability of the endothelial cell layer, thus promoting the entry of amyloid-β peptides from the serum into the central nervous system, where they bind to neurons. We also found that hydrocephalous occurred in some of the hSlit2 transgenic mice. In addition, we observed a larger lateral ventricle area and higher ventricle pressure in the transgenic mice. A comparison of the choroids plexus, where CSF is secreted, revealed that transgenic mice have changes in the structure of the choroids plexus, including more microvessels, dilated vessels, and gaps between epithelial cells and endothelial cells. Thus, Slit2 could bind to its receptor Robo1 on endothelial cells and affect the junction through signaling in the cytoplasm. It has been reported that Slit2 modifies the activity of cytoplasmic GTP enzymes, which affect tight junction structures by regulating adherence proteins and cell scaffold proteins[37]. However, the signaling pathway by which Slit2 affects cell-cell adhesion structures requires further investigation. Because adherent structures consist of a large number of proteins, it is unclear which proteins are affected by Slit2. For example, which changes in protein expression or function are the direct results of Slit2 signaling, and which

Figure 5. hSlit2 expression changes blood vessel permeability at sites of cell-cell adhesion. (A) Permeability of C57 and hSlit2 mouse brains. The data represent the mean results from seven mice; bars, SEM. *P<0.01. (B) Measurement of the ICH area in brain longitudinal section slides from C57 and hSlit2 mice. The data represent the mean results from six mice; bars, SEM. *P<0.01. (C and D) Longitudinal section slides of mouse brains in which ICH was induced. The ICH area was bigger in Slit2 mice (D) than in C57 mice (C). (E and F) Immunofluorescence of HUVECs with anti VE-cadherin antibody. Cell-cell adhesion was different in hSlit2 stimulated HUVECs (F) compared with cells not stimulated with hSlit2 (E). (G) hSlit2 increases the permeability of HUVECs in in vitro permeability assays. The Robo blocking antibody R5 could block this effect.
protein changes are the result of subsequent disturbances of junction structures? These are all important questions that need to be answered to understand the molecular mechanism by which Slit2 alters the permeability of barriers.

In the brains of transgenic mice of 6-week-old, we observed binding of the amyloid-β peptide to neurons. This suggested that Slit2 promotes the entry of amyloid-β peptides from the serum into the central nervous system, where they then bind to neurons. The binding of amyloid-β peptides to neurons is a phenotype of the early stage of Alzheimer’s disease. However, whether Slit2 is over-expressed in patients with Alzheimer’s disease and the relationship between Slit2 and Alzheimer’s disease need to be further investigated and will be the focus of future work in our lab.

Slit2 improves the permeability of the blood-brain barrier and thus may have some medical application for the delivery of drugs to the central nervous system, a problem that has puzzled many researchers because drugs targeting the central nervous system often cannot penetrate the blood-brain barrier at sufficient therapeutic doses. In our study, we found that Slit2 promotes the entry of a fluorescent-labeled peptide into the central nervous system. This result suggests that Slit2 may promote the penetration of large-molecule drugs from the peripheral circulation into the central nervous system by increasing the permeability of the blood-brain barrier. Whether manipulation of Slit2 levels or activity can be applied to the field of central nervous system drug delivery requires further investigation, which we plan on pursuing in the future.

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Author contribution
Hai-xiong HAN and Jian-guo GENG designed the research. Hai-xiong HAN performed the research, analyzed the data, and wrote the paper.

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Figure 7. Structure of the aqueduct and SAS and expression of VEGF in transgenic and C57 mice. (A, B and C, D) Structure of the aqueduct in Slit2 mice (A, C) and C57 mice (B, D). No difference was observed in the aqueduct between Slit2 and C57 mice. (E, F and G, H) Structure of the SAS in Slit2 mice (E, G) and C57 mice (F, H). No difference was observed in the SAS between Slit2 and C57 mice. (I) VEGF concentrations in the blood of C57 and Slit2 mice. Serum VEGF levels of C57 and Slit2 mice were detected with a VEGF ELISA kit (Quantikine Immunoassay Lot: 234711). The data represent the mean results for groups of eight mice were detected with a VEGF ELISA kit. (J) Q-PCR analysis of VEGF mRNA extracted from C57 and hSlit2 mice. VEGF expression was not increased by hSlit2 in mouse brains.

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