The S1 protein of SARS-CoV-2 crosses the blood–brain barrier in mice

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It is unclear whether severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), which causes coronavirus disease 2019 (COVID-19) pandemic, can enter the brain. Severe acute respiratory syndrome coronavirus 2 binds to cells via the S1 subunit of its spike protein. We show that intravenously injected radiolabeled S1 (I-S1) readily crossed the blood–brain barrier in male mice, was taken up by brain regions and entered the parenchymal brain space. I-S1 was also taken up by the lung, spleen, kidney and liver. Intranasally administered I-S1 also entered the brain, although at levels roughly ten times lower than after intravenous administration. APOE genotype and sex did not affect whole-brain I-S1 uptake but had variable effects on uptake by the olfactory bulb, liver, spleen and kidney. I-S1 uptake in the hippocampus and olfactory bulb was reduced by lipopolysaccharide-induced inflammation. Mechanistic studies indicated that I-S1 crosses the blood–brain barrier by adsorptive transcytosis and that murine angiotensin-converting enzyme 2 is involved in brain and lung uptake, but not in kidney, liver or spleen uptake.

SARS-CoV-2 is responsible for the coronavirus disease 2019 (COVID-19) pandemic. In addition to pneumonia and acute respiratory distress, COVID-19 is associated with a host of symptoms that relate to the CNS, including loss of taste and smell, headaches, twitching, seizures, confusion, vision impairment, nerve pain, dizziness, impaired consciousness, nausea and vomiting, hemiplegia, ataxia, stroke and cerebral hemorrhage. It has been postulated that some of the symptoms of COVID-19 may be due to direct actions of the virus on the CNS; for example, respiratory symptoms could be in part due to SARS-CoV-2 invading the respiratory centers of the brain. Encephalitis has also been reported in COVID-19, and could be a result of virus or viral proteins having entered the brain. SARS-CoV-2 mRNA has been recovered from the cerebrospinal fluid, suggesting it can cross the blood–brain barrier (BBB). Other coronaviruses, including the closely related SARS virus that caused the 2003–2004 outbreak, are able to cross the BBB, and SARS-CoV-2 can infect neurons in a BrainSphere model. However, SARS-CoV-2 could induce changes in the CNS without directly crossing the BBB, as COVID-19 is associated with a cytokine storm, and many cytokines cross the BBB to affect CNS function.

Here we assess whether one viral protein of SARS-CoV-2, the spike 1 protein (S1), can cross the BBB. This question is important and clinically relevant for two reasons. First, some proteins shed from viruses have been shown to cross the BBB, inducing neuroinflammation and otherwise impairing CNS functions. Second, the viral protein that binds to cells can be used to model the activity of the virus; in other words, if the viral binding protein crosses the BBB, it is likely that protein enables the virus to cross the BBB as well. S1 is the binding protein for SARS-CoV-2 (ref. 2); it binds to angiotensin-converting enzyme 2 (ACE2) and probably other proteins as well.

In this study, we show that I-S1 readily crossed the murine BBB, entered the parenchymal tissue of the brain and, to a lesser degree, was sequestered by brain endothelial cells and associated with the brain capillary glyocalyx. We describe I-S1 rate of entry into the brain after intravenous (i.v.) and intranasal administration, determine its uptake in 11 different brain regions, examine the effect of inflammation, APOE genotype and sex on I-S1 transport, and compare I-S1 uptake in the brain to the uptake in the liver, kidney, spleen and lung. Based on experiments with the glycoprotein WGA, we found that brain entry of I-S1 likely involves the vesicular-dependent mechanism of adsorptive transcytosis.

Results

I-S1 protein is transported across the mouse blood–brain barrier. We obtained S1 proteins from two commercial sources: RayBiotech and AMSBIO. The S1 proteins were radiolabeled in-house, and verified to be intact after labeling by autoradiography gels (Extended Data Fig. 1 and Supplementary Fig. 1). We determined whether intravenously injected I-S1 could cross the BBB in mice, by measuring its blood-to-brain influx constant (K) using multiple-time regression analysis (MTRA). MTRA plots the tissue/serum ratios for I-S1 against exposure time, which is a measure of time that has been corrected for the clearance of I-S1 from blood. The slope of the linear portion of this plot measures K, that is, the unidirectional influx constant for I-S1.

We co-injected 99mTc-labeled albumin (T-Alb) along with the I-S1. T-Alb crosses the intact BBB poorly and so can be used to measure the vascular space of the brain. Any brain/serum ratios for I-S1 that exceed the brain/serum ratios of T-Alb therefore represent extravascular I-S1; that is, material which has crossed the BBB. T-Alb can also be used to measure the leakiness of peripheral tissue beds and of a BBB that has been disrupted by disease or inflammation.
I-S1 is cleared from blood and taken up by peripheral tissues. We determined the clearance of intravenously injected I-S1 from RayBiotech from blood and its uptake by brain and other tissues (Fig. 2). Clearance of I-S1 from blood was linear for the first 10 min, with a half-time clearance of about 6.6 min (Fig. 2a), followed by a plateau. In Fig. 2 and subsequent figures, tissue/serum ratios for T-Alb were subtracted from the tissue/serum ratios for I-S1. These ‘delta’ tissue/serum ratios were thus corrected for the I-S1 in the vascular space and for any I-S1 that would have entered tissue because of a leaky capillary bed. The resulting delta values thus represent specific uptake of I-S1 by the tissues.

All tissues showed uptake of I-S1 (Fig. 2b–f). Spleen and liver uptake was nonlinear, suggesting that their tissue beds were coming into equilibrium with blood. Most substances in blood are cleared by kidney or liver; the much higher I-S1 uptake in liver compared to kidney suggests that I-S1 is cleared from blood predominantly by the liver. To determine if there were regional differences in I-S1 uptake within the brain, we collected the olfactory bulb and disected the whole brain into ten regions (Extended Data Fig. 2). We found that I-S1 entered all brain regions, with no statistically significant differences among them.

We similarly determined the clearance of I-S1 (AMSBIO) from blood and its uptake by brain and other tissues (Fig. 3). The results were similar to those obtained with I-S1 derived from RayBiotech (Fig. 2), but there were a few differences. The AMSBIO-derived I-S1 was cleared from blood faster (half-life of 3.6 min), and the rate of liver uptake was nearly five times faster than liver uptake of RayBiotech-derived I-S1 (P < 0.0001). Together, these results show that I-S1 is cleared from blood primarily by the liver and that S1 has access to other tissues relevant to COVID-19, including kidney, lung and spleen.

I-S1 is stable in brain and blood. We measured the stability of intravenously injected I-S1 from RayBiotech in blood and brain through acid precipitation (Supplementary Table 1). I-S1 added to tissues (processing controls) ex vivo showed little degradation. Radioactivity recovered from brain 10 min after i.v. injection and serum 10 and 30 min after the i.v. injection of I-S1 was stable, with most of the radioactivity precipitated by acid. Radioactivity recovered from brain 30 min after the i.v. injection of I-S1 was mostly degraded. This indicates that I-S1 enters the BBB intact but is eventually degraded in the brain, although it is not clear whether this involves cleavage of the radioactive iodine from I-S1 and/or whether S1 protein itself is degraded.

Most of the I-S1 taken up by the capillary bed enters the brain parenchyma. In rare cases, substances that seem to cross the BBB are actually sequestered by the capillary bed, which prevents them from entering the brain parenchymal/interstitial fluid space. Therefore, we next assessed whether intravenously injected I-S1 fully crosses the capillary wall to enter brain parenchyma. For this, we used a modified version of the ‘capillary depletion’ method that also enabled us to measure the amount of intravenously injected I-S1 that reversibly adheres to the luminal side of the capillary bed. We found that binding of I-S1 to the luminal side remained static over time, whereas there was a small increase over time in the amount of I-S1 retained by brain capillaries (Extended Data Fig. 3). The largest change over time was in the amount of I-S1 entering the parenchyma. These results show that by 30 min, over 50% of I-S1 had crossed the capillary wall fully to enter into the brain parenchymal and interstitial fluid spaces.

WGA enhances I-S1 uptake in brain and peripheral tissues. Wheatgerm agglutinin (WGA) is a plant lectin that crosses the BBB through adsorptive transcytosis—a process by which proteins or peptides bind to glycoproteins of the luminal surface of endothelial cells, are internalized into vesicles, and then transported across the membrane. For WGA, adsorptive transcytosis occurs when WGA binds to cell-surface glycoproteins containing siaic acid and N-acetylgalactosamine. Many viral proteins also bind to glycoproteins containing siaic acid and N-acetylgalactosamine, and therefore, when WGA is co-injected with such viral proteins, it enhances their transport across the BBB and uptake in peripheral tissues15. Here, we found that uptake of intravenously injected I-S1 (RayBiotech or AMSBIO) into brain, lung, spleen and kidney was increased when WGA was included in the injection (Fig. 4b–e). WGA co-injection also increased clearance of I-S1 (RayBiotech), but not I-S1 (AMSBIO), as shown by a decrease in I-S1 in blood (Fig. 4a). WGA co-injection decreased uptake of I-S1 (AMSBIO) but not I-S1 (RayBiotech) in the liver (Fig. 4f), although it did decrease I-S1 (RayBiotech) uptake in liver in another experiment (see below and Fig. 4h).

These results strongly suggest that I-S1 crosses the BBB and peripheral tissue beds through the mechanism of adsorptive
transcytosis, binding to cell-surface glycoproteins that contain sialic acid or N-acetylgalacosamine.

Heparin blocks uptake of I-S1 in liver but not in brain. Some viruses use heparan sulfate on the cell membrane as a receptor\cite{25}. Heparin can block virus uptake in tissues, presumably by binding to viral proteins and thereby blocking these viral proteins from binding to the heparan sulfate on the cell membranes\cite{25, 26}. However, we found that heparin co-injected with I-S1 did not affect uptake of I-S1 (RayBiotech) in brain, lung, spleen or kidney and did not influence the effect of WGA co-injection on I-S1 uptake in these tissues (data not shown). Heparin co-injected with I-S1 did block uptake of I-S1 in liver (Fig. 4h), which likely explained the observed decreased clearance of I-S1 from blood (as evidenced by the increase in I-S1 levels in serum; Fig. 4g), but it did not block the effect of WGA on I-S1 uptake in liver. These results indicate that I-S1 uses heparin-sensitive sites to bind to the capillary bed of the liver, but not to bind to the capillary beds of brain, spleen, lung or kidney.

I-S1 uptake in brain was not blocked by unlabeled S1. We next determined whether transport of intravenously injected I-S1 (RayBiotech) into the brain could be blocked by including in the injection an excess of unlabeled S1 protein (AMSBIO). Unlabeled S1 did not affect brain/serum ratios for delta I-S1 at any of the tested doses (Fig. 5a), although it did enhance uptake of I-S1 in lung (Fig. 5c). This indicates that the binding site for I-S1 in brain tissue is not...
Fig. 3 | I-S1 (AMSBIO) is cleared from blood and taken up by peripheral tissues. a, Clearance of I-S1 from AMSBIO from blood. Data were fitted to a one-phase exponential decay model: \( y = 1.05 (e^{-0.39t}) + 0.880 \) (half-life of 1.74 min; \( n = 11 \) mice; \( r^2 = 0.867 \)). Clearance from blood was linear for the first 10 min, with a half-time clearance of 3.6 min. b–f, the y axes show the delta tissue/serum ratios for I-S1, corrected for vascular space and nonspecific leakage (Methods). b, The I-S1 \( K_i \); value for whole brain was significantly non-zero (\( P < 0.0001 \)), indicating tissue uptake (\( n = 11 \) mice). c–f, The I-S1 \( K_i \); values for different tissues calculated based on the linear portions of their curves. c, The I-S1 \( K_i \); for lung was significantly non-zero (\( P < 0.0001; n = 11 \) mice). d, The I-S1 \( K_i \); value for spleen was significantly non-zero (\( P = 0.002; n = 11 \) mice). e, The I-S1 \( K_i \); value for kidney was significantly non-zero (\( P = 0.0004; n = 11 \) mice). f, The I-S1 \( K_i \); for liver was significantly non-zero (\( P = 0.0005; n = 8 \) mice; \( n = 3 \) mice (filled circles) were excluded from regression analysis due to nonlinearity). All error terms are the s.e.m.

ACE2 likely mediates I-S1 uptake in brain and lung but not other tissues. The ability of SARS-CoV-2 to enter cells is thought to depend on the S protein binding to the membrane-bound enzyme (and glycoprotein) ACE2. To assess whether ACE2 has a role in I-S1 uptake in brain and peripheral tissues, we intravenously co-injected human ACE2 or the ACE2 substrates Ang II and ghrelin with I-S1 (RayBiotech). This did not affect blood levels of intravenously injected I-S1 or T-Alb (data not shown), indicating that these proteins did not affect the volume of distribution or clearance of either S1 or albumin. Co-injection of ACE2 increased kidney levels of intravenously injected T-Alb: \( F(4,36) = 2.63, P = 0.0505; \) vehicle: 119 ± 15.6 \( \mu \)g \( g^{-1} \) (\( n = 8 \) mice); ACE2: 151 ± 15.6 \( \mu \)g \( g^{-1} \) (\( n = 8 \) mice), \( P = 0.02 \), possibly indicating that ACE2 affects renal clearance of albumin. Only ACE2 co-injection enhanced I-S1 uptake in brain (Fig. 5b). Uptake in lung was increased by co-injection of S1, ghrelin or ACE2, but not angiotensin II (Fig. 5c), whereas uptake in liver, kidney or spleen was not altered by co-injection of any of these substances (data not shown). These results suggest that ACE2 is involved in S1 uptake in lung and probably brain, but not in I-S1 uptake in spleen, liver or kidney.

Inflammation increases I-S1 uptake in brain and lung. Since SARS-CoV-2 infection induces an inflammatory state, we next
Fig. 4 | WGA enhances I-S1 uptake by brain, lung, spleen and kidney, while heparin blocks uptake by liver. 

**a-f.** Serum levels and tissue uptake of I-S1 from either RayBiotech or AMSBIO 10 min after i.v. injection in which WGA, a stimulator of adsorptive transcytosis, was included in the injections to half of the mice. Serum I-S1 levels expressed as the percentage of the injected dose per ml of serum. There was a main effect of S1 source (F(1,25) = 49.79, P < 0.0001) and treatment (F(1,25) = 20.57, P < 0.0001), as well as a significant effect of their interaction (F(1,25) = 40.93, P < 0.0001). WGA treatment decreased I-S1 (RayBiotech) levels in serum (P < 0.0001), indicating increased clearance from blood. Brain I-S1 levels expressed as delta brain/serum ratios, showing significant effects (all P < 0.0001) of S1 source (F(1,25) = 92.98), treatment (F(1,25) = 252.2) and their interaction (F(1,25) = 91.41). These results show that WGA stimulated I-S1 uptake from both RayBiotech and AMSBIO and that WGA stimulated uptake of I-S1 from AMSBIO more than from RayBiotech. Lung I-S1 levels show an effect of treatment (F(1,25) = 204.6, P < 0.0001). Spleen I-S1 levels show effects of S1 source (F(1,25) = 43.06, P < 0.0001) and treatment (F(1,25) = 30.42, P < 0.0001) and a trend for interaction (F(1,25) = 3.657, P = 0.08). Kidney I-S1 levels show effects of source (F(1,25) = 33.45, P < 0.0001) and treatment (F(1,25) = 40.77, P < 0.0001). Liver I-S1 levels show effects of source (F(1,25) = 55.87), treatment (F(1,25) = 52.16) and their interaction (F(1,25) = 32.08), all at P < 0.0001. For **a-f,** data were analyzed using two-way ANOVA with S1 source (RayBiotech versus AMSBIO) and treatment (with or without WGA) as main factors, followed by Tukey’s multiple-comparisons tests; data are the mean ± s.d. **P < 0.05, ***P < 0.01 and ****P < 0.001, indicating comparisons of treatments within I-S1 group (RayBiotech or AMSBIO); **P < 0.05, ***P < 0.01 and ****P < 0.001, indicating comparisons of I-S1 groups within treatments. n = 7 per group, except for RayBiotech S1-WGA mice (n = 8).**

**g.** Serum levels and liver uptake of I-S1 from RayBiotech 10 min after i.v. injection with or without co-injection of WGA, heparin or both. Heparin increased serum I-S1 (% inj ml–1) ratios, showing significant effects (all P < 0.0001) of S1 source (F(1,25) = 30.08) and their interaction (F(1,25) = 20.57, P < 0.0001), indicating comparisons of I-S1 groups within treatments. n = 7 per group, except for RayBiotech S1-WGA mice (n = 8). **h.** In comparison to vehicle, uptake of I-S1 was decreased by heparin (P = 0.0015), WGA (P < 0.0001) and WGA + heparin (P < 0.001). Liver uptake of I-S1 was lower after WGA + heparin versus heparin alone (P = 0.0034) but not versus WGA alone. *P < 0.05, **P < 0.01 and ***P < 0.001. For **g** and **h,** data were analyzed by one-way ANOVA and Sidak’s multiple-comparisons test, with n = 10 for each group, with the exception of liver vehicle in **h,** where one outlier was identified using a Grubbs test (77.6 μl g–1) and excluded, leaving n = 9 mice.
determined whether an inflammatory state—in this case, induced by lipopolysaccharide (LPS) injection—affects uptake of intravenously injected I-S1 (RayBiotech) in brain and peripheral tissues. Mice received a 3 mg kg⁻¹ injection of LPS derived from *Salmonella typhimurium* (Sigma) at *t*=0 and again at 6 and 24 h later, and received an i.v. I-S1 plus T-Alb injection 28 h after the first LPS injection; this LPS regimen can disrupt the BBB, increase its permeability to viruses and viral proteins, and increase the blood level of many of the cytokines found to be elevated in the cytokine storm associated with COVID-19 (refs. 52-56). 

We found that LPS did not affect T-Alb blood levels, indicating that inflammation did not induce volume contraction of the vascular space nor leakiness of the peripheral capillary beds in these mice. Mice that had received LPS had higher I-S1 serum levels (Fig. 6a), indicating a decrease in clearance from blood (likely because of the decreased I-S1 uptake in liver observed in these mice; Fig. 6b), as well as higher T-Alb brain/serum ratios, indicating BBB disruption (control: 8.52 ± 0.19 μ g⁻¹ (n = 10); LPS: 12.2 ± 0.68 μ g⁻¹ (n = 8), *t*= 5.74, *P* < 0.0001). These mice also had higher I-S1 brain/serum ratios (that is, ratios not corrected for I-S1 in the vascular space and for any I-S1 brain entry due to leakage), indicating increased I-S1 passage across the BBB (control: 12.06 ± 0.26 μ g⁻¹ (n = 10); LPS: 14.9 ± 0.64 μ g⁻¹ (n = 8), *t*= 4.38, *P*= 0.0005), although their delta brain/serum ratios for I-S1 (that is, corrected ratios) were not different from controls (except in the olfactory bulb) (Fig. 6c). The only peripheral tissue showing an increased uptake of I-S1 after receiving LPS was the lung, where uptake was increased by 101% (Fig. 6b). These results show that inflammation could increase S1 toxicity for lung tissue by increasing its uptake. The results also show that, in mice, inflammation can increase entry of intravenously injected I-S1 into brain, but this is likely due to BBB disruption rather than by enhancing absorptive transcytosis.
I-S1 enters mouse brain and blood after nasal administration. Some viruses can enter the brain via the olfactory nerve, a cranial nerve with projections through the cribriform plate. Other viruses may cross the blood-brain barrier from the nasal compartment. It is unclear whether SARS-CoV-2 in the nose can enter brain by either of these routes. We therefore assessed the ability of intranasally administered I-S1 to enter the brain. Following an injection of a 1-µl vehicle solution containing I-S1 to each naris at the level of the cribriform plate, I-S1 was detected in all brain regions (Fig. 7a,b). I-S1 levels in whole brain, frontal cortex, cerebellum, midbrain and pons-medulla were higher at 30 min compared to 10 min after administration. I-S1 distribution in the brain was mostly homogeneous, although levels in the olfactory bulb and hypothalamus were higher than in other brain regions at 10 min and also in the olfactory bulb at 30 min after administration. Whole-brain I-S1 levels expressed as the percentage of the administered dose per 1 ml (% inj ml⁻¹) of serum, were increased 4 h after LPS injection versus control (t = 3.66, df = 16, P = 0.0021, two-tailed t-test), indicating that LPS decreased clearance of I-S1 from blood. LPS had no significant effect on the serum levels of I-S1 (t = 0.3739, df = 16, P = 0.7134), indicating a selective effect of LPS on I-S1 clearance. Tissue levels of I-S1, expressed as tissue/serum ratios corrected (delta brain/tissue ratios) for vascular space and leakage by subtracting the tissue/serum ratios for T-Alb. LPS increased I-S1 uptake by lung tissue (t = 4.58, df = 16, P = 0.0003) and decreased I-S1 uptake by spleen (t = 6.39, df = 16, P < 0.0001) and liver (t = 4.60, df = 16, P = 0.0003); the effect of LPS on I-S1 uptake by kidney was not significant (P = 0.057; all two-tailed t-tests). The decreased I-S1 uptake by liver tissue likely contributes to the decrease in I-S1 clearance from blood seen in a, c. Levels of I-S1 in brain areas, expressed as delta brain/serum ratios. Two-way ANOVA showed a significant main effect of brain region (F(11, 190) = 2.72, P = 0.0028), accounting for 12.5% of the variability, and a main effect of LPS treatment (F(11, 190) = 10.7, P = 0.0012), accounting for 4.5% of the variability; there was no statistically significant effect for interaction. Sidak’s multiple-comparisons test showed that LPS treatment decreased I-S1 uptake by the olfactory bulb (P = 0.026) but not other brain regions. *P < 0.05, **P < 0.01, ***P < 0.001 and t = 0.1 > P > 0.05. FCtx, frontal cortex; PCtx, parietal cortex; OCtx, occipital cortex. Data are the mean ± s.d. Control groups (a): n = 10 mice; vehicle group (b and c): n = 10 mice; LPS group (a–c): n = 10 mice.
Fig. 7 | I-S1 enters the brain after intranasal administration. a, b. Brain uptake of I-S1 (RayBiotech) delivered intranasally (1 μl per naris), expressed as the percentage administered dose taken up per gram of each brain region (% inj g⁻¹): a. At 10 min after intranasal administration, I-S1 was detected in all dissected brain regions. One-way ANOVA showed a significant effect of brain region (F(11,48) = 3.777, P = 0.0006), with significantly higher uptake in the olfactory bulb (P = 0.0031) and hypothalamus (P = 0.0041) compared to the whole brain (Dunnett’s multiple-comparisons test). Whole-brain uptake after intranasal administration was 0.023 ± 0.0008 % inj g⁻¹; for visual comparison, we also show the whole-brain percentage dose per gram 10 min after i.v. injection as calculated from the control values for the LPS experiment in Fig. 6 (filled circles, 0.22 ± 0.016 % inj g⁻¹); n = 5 mice. b. At 30 min after intranasal administration, I-S1 was detected in all dissected brain regions. One-way ANOVA showed a significant effect of brain region (F(11,48) = 4.33, P = 0.0002), with significantly more uptake in the olfactory bulb compared to whole brain (P = 0.0023, Dunnett’s multiple-comparisons test). A two-tailed t-test showed significant increases between 10 and 30 min for the whole brain (P = 0.0143, t = 3.116, df = 8), frontal cortex (P = 0.0391, t = 2.464, df = 8), cerebellum (P = 0.0003, t = 6.226, df = 8), midbrain (P = 0.0098, t = 3.370, df = 8) and pons (P = 0.0232, t = 2.801, df = 8) but not the other regions; n = 5 mice. *P < 0.05, **P < 0.01 and ***P < 0.001.

Effects of sex and APOE genotype on I-S1 uptake. Male sex and APOE4 genotype in comparison to the APOE3 genotype are risk factors for both contracting and having a poor outcome from COVID-19 (refs. 31–33). Therefore, we determined uptake of intravenously injected I-S1 (RayBiotech) or T-Alb in male and female mice expressing human APOE3 or APOE4 under the expression of the mouse ApoE promoter. APOE genotype and sex did not affect T-Alb brain/serum ratios (Fig. 8a). Two-way ANOVA showed an effect of sex on I-S1 uptake in the olfactory bulb and spleen and an effect of APOE genotype on I-S1 uptake in liver, spleen and kidney (Fig. 8b–h). Multiple-comparisons tests showed that, compared to the other three groups, male mice expressing human APOE3 had the fastest I-S1 uptake of I-S1 in the olfactory bulb, liver and kidney, whereas compared to the other three groups, female mice expressing human APOE3 had the fastest I-S1 uptake in spleen. These results suggest that enhanced uptake of I-S1 by some tissues could contribute to the increased risk of COVID-19 in males versus females, but not to the risk associated with the APOE4 genotype.

I-S1 transport across an in vitro model of the BBB. To determine whether the I-S1 crosses endothelial-like cells in an in vitro model of the human BBB, we compared transport of I-S1 versus T-Alb across monolayers of human induced pluripotent stem cell (iPSC)-derived brain endothelial-like cells (iBECs) seeded on Transwells. Three independent experiments showed that rate of passage of I-S1 (RayBiotech) in the luminal-to-abluminal direction was not statistically significant when compared to that of T-Alb (Extended Data Fig. 4a). One experiment indicated that unlabeled S1 did not inhibit I-S1 (RayBiotech) passage (Extended Data Fig. 4b). WGA also had no effect on RayBiotech I-S1 transport (Extended Data Fig. 4c). I-S1 (RayBiotech) had a significantly higher permeability coefficient than I-S1 (AMSBIO I-S1) in vitro (Extended Data Fig. 4d); however, this difference was not significant after correcting for T-Alb transport. In summary, these results suggest that iBECs may be more permeable to I-S1 than to T-Alb, but the difference is too small to support mechanistic studies using this model.

Discussion
The results from this study show that I-S1 from two different commercial sources readily crosses the mouse BBB, at least when injected intravenously. I-S1 was taken up by all 11 brain regions examined. Such widespread entry into brain of I-S1 could explain the diverse effects of S1 and/or SARS-CoV-2 such as encephalitis, respiratory difficulties and anosmia1,4. S1 is the SARS-CoV-2 protein that initially binds to cell-surface receptors, setting the stage for viral internalization. For transport across the BBB, viral binding proteins often behave similarly to the virus itself. For example, interactions (including binding and transport) between the HIV-1 glycoprotein gp120 and the BBB are similar to those for the complete virus16,20. Additionally, many if not most viral proteins themselves can be biologically highly active; for example, gp120 is highly toxic11–17. Coronavirus spike proteins are often cleared from the virus by host cell proteases. Once cleaved, coronavirus spike S1 and S2 subunits are not held covalently by disulphide bonds and so S1 could be shed from virions14. It is possible that during infection by SARS-CoV-2, shed S1 is available to cross the BBB, triggering responses in the brain itself, without necessarily involving crossing of intact virus particles. Thus, determining whether S1 crosses the BBB is important for understanding whether SARS-CoV-2 and S1 itself could induce responses in the brain.

Our method of studying S1 pharmacokinetics has many advantages over the more traditional approach of determining viral uptake and distribution that is based on virus recovery. Radioactive labeling allows S1 to be detected at very low levels and quantification of the rates of uptake for brain and other tissues. Factors that might affect viral protein uptake can be manipulated experimentally in healthy, rather than infected, animals. Recovery of I-S1 from a tissue reflects only factors related to permeability, whereas recovery from infected mice also reflects other factors, such as rate of virus replication in that tissue.

A crucial question which we partially answered here was: what receptor does I-S1 use to enter brain and other tissues? Based on
experience with SARS, it is assumed that SARS-CoV-2 will bind to human ACE2, but not murine ACE2. SARS can infect wild-type (WT) mice, but it doesn't produce severe symptoms and death, except in transgenic mice overexpressing human ACE2 (ref. 35), although this could also simply be because these transgenic mice express 8–12 times more ACE2 than WT mice. The mice used here only expressed murine receptors, so our findings suggest that the assumption that ACE2 must be the human protein is incorrect and demonstrate that WT mice can be used in kinetics studies of S1 and probably SARS-CoV-2.

We did find reasonably strong evidence that murine ACE2 is involved in I-S1 uptake in lung tissue, as co-injection of I-S1 with soluble ACE2 and ACE2 substrates increased the uptake of I-S1 (this perhaps surprising observation is discussed further below). The evidence for ACE2 involvement in brain I-S1 uptake is weaker than that for lung, as here, uptake was affected by co-injection with soluble ACE2, but not by ACE2 substrates. The finding that I-S1 uptake in the kidney, liver and spleen was unaffected by soluble ACE2 or by ACE2 substrates indicate that receptors other than, or in addition to, ACE2 are involved in uptake of I-S1 in some tissues.

That S1 and even SARS-CoV-2 would use more than one receptor is not surprising when one considers that many viruses use multiple receptors. For example, HIV-1 uses the CD4 and mannose-6-phosphate receptors, and the rabies virus uses the acetylcholine receptor, a nerve growth factor receptor and the neural cell adhesion molecule to enter cells36–38. Receptors (besides ACE2) that can bind or are predicted to bind SARS-CoV-2 based on modeling include basigin, cyclophilins, dipetidyl peptidase-4 (refs. 39,40) and GRP78 (ref. 24). One reason that a virus can use such a diversity of receptors is that viruses bind with less specificity than do endogenous receptor ligands. The binding sites for viral proteins are often highly charged regions on the cell-membrane glycoprotein owing to high concentrations of sialic acid, N-acetylgalactosamine or heparan sulfate. Coronaviruses in general bind to glycoproteins high in sialic acid41. Pioneering work showed that WGA binding to BBB regions rich in sialic acid or N-acetylgalactosamine resulted in the transportation of WGA across the BBB through the mechanism of adsorptive transcytosis24. WGA coadministered with a weaker inducer of adsorptive transcytosis will often increase rather than block the BBB penetration of the weaker inducer41. In the current study, the ability of WGA to increase the I-S1 uptake in brain tissue suggests that S1 crosses the BBB through adsorptive transcytosis.

Because the spike protein of SARS-CoV-2 is more highly charged than the spike protein of SARS, it has been suggested that it may bind to a larger number of receptors42. Some viruses bind receptors rich in heparan sulfate; uptake of those viruses is inhibited by heparin43. We showed that I-S1 uptake in liver was inhibited by heparin, but uptake in brain and other peripheral tissues was not. These results show that S1 uses heparan sulfate to bind to liver but not to other tissues. We conclude that a number of receptors are likely involved in S1 uptake; which receptor is most important varies from tissue to tissue. It will be important to identify the membrane-bound glycoproteins that serve as receptors for SARS-CoV-2.

It is unclear why, in our study, co-injection with ACE2 or ACE2 substrates enhanced rather than inhibited uptake of intravenously injected I-S1, but there are some possible explanations. Since S1 does not bind to the ACE2 catalytic site44,45, traditional ligand–receptor dose-dependent inhibition kinetics may not occur. The ACE2 we co-injected with I-S1 may have bound circulating Ang II that would normally have competed with I-S1 for binding to membrane-bound ACE2. In addition, S1 is attached to SARS-CoV-2 as a homotrimer46, but we studied monomeric S1; it is possible that co-injection of ACE2 ligands altered ACE2 conformation in such a way that it facilitated binding of the S1 monomer.

Risk factors for both contracting COVID-19 and having a poor outcome include male individuals who are positive for the ApoE4 allele in comparison to the ApoE3 allele31–33, whereas cytokine storm is a characteristic of severe disease46. We found that the influence of sex, ApoE genotype and inflammatory state on I-S1 uptake varied among the tissues. Sex and human ApoE status in mice did not affect uptake of intravenously injected I-S1 in whole brain or lung, but did affect its uptake in olfactory bulb, liver, spleen and kidney, with higher uptake in males. This suggests that some of the risk of poor outcome for males may be related to the degree to which their tissues have an enhanced uptake of S1 or SARS-CoV-2. However, ApoE3, not ApoE4, was associated with higher uptake rates of I-S1 by the olfactory bulb, liver, spleen and kidney, suggesting that the risk associated with ApoE status is unlikely to be due to increased S1 or SARS-CoV-2 tissue uptake.

Inflammation induced by LPS injection increased the amount of intravenously injected I-S1 entering the brain, but this increase was likely due to BBB disruption and not due to enhancement of adsorptive transcytosis; indeed, uptake of S1 after correction for BBB disruption was actually lower in one brain region, the olfactory bulb. LPS-exposed mice had higher I-S1 uptake in lung but lower uptake in spleen and liver; the latter likely explains why these mice had reduced I-S1 clearance from blood. Notably, this decrease in clearance from blood observed in mice in an inflammatory state suggests that all tissues will be exposed to higher S1 levels than in the noninflammatory state.

Fig. 8 | Sex and ApoE genotype influence on I-S1 uptake. a–h. Serum levels and tissue uptake of I-S1 (RayBiotech) and brain uptake of T-Alb following i.v. injection in male and female mice that were homozygous for human APOE3 (E3) or APOE4 (E4). For a, MTRA was performed to calculate the K, and V, for T-Alb. In c–h, the y axis shows delta tissue/tissue ratios for I-S1, which were corrected for vascular space and nonspecific leakage (Methods). The slopes of the regression lines that result from MTRA measures the rate of influx (Kd) and to E3 males compared to E3 females (P = 0.0046), accounting for 25% of variability, with faster uptake in E3 males compared to E3 females (P = 0.0057) and to E4 females (P = 0.0015). APOE genotype had a trend-level effect on uptake in the olfactory bulb (F(1,29) = 3.838, P = 0.06). f, APOE genotype had a significant main effect on I-S1 uptake in the liver (F(1,24) = 28.44, P < 0.0001), accounting for 50% of the variability, with a lower uptake rate for E4 females compared to E3 females (P = 0.0057) and to E3 males (P = 0.0001), and with a lower uptake rate for E4 males versus E3 males (P = 0.0039).

There was a significant main effect of sex on I-S1 uptake in the spleen (P = 0.0004), accounting for 20% of the variability, a main effect of ApoE genotype (P < 0.0001), accounting for 43% of variability, and a significant effect of their interaction (P = 0.0029), accounting for 13% of variability. Multiple comparisons showed faster uptake rates for E3 females compared to E4 females (P < 0.0001) to E3 males (P = 0.0003) and to E4 males (P < 0.0001). h, APOE genotype had a significant effect on I-S1 uptake in the kidney (P = 0.0046), accounting for 25% of variability, with faster uptake in E3 males compared to E4 males (P = 0.0089) and to E4 females (P = 0.012). The number of mice assigned each group was 11 for E3 females and 10 for the other three groups. However, the number of mice used to calculate the regression lines ranged from 6 to 11 per group, because deviation from linearity was observed that satisfied exclusion criteria (Methods). The n values per group (including the excluded outliers) is shown in Supplementary Table 2.
A lethal infection can occur after intranasal administration of SARS-CoV-2. It has been postulated that nasal virus spreads to the lung and from there to blood and brain, but others suggest that SARS-CoV-2 in the nares could spread to the brain through the olfactory nerve, as do many other viruses. Although our findings show that intranasally administered I-S1 can enter mouse brain tissue, they highlight the BBB as the major route for I-S1 entry into the brain. Moreover, a very small amount (0.66% bioavailability after intranasal administration) of I-S1 was found in blood, suggesting poor nasal-to-blood transfer. However, our studies were designed to assess the ability of I-S1 to enter the brain through the olfactory nerve and not to assess its ability to enter the blood via the nasal vasculature. Nevertheless, our results favor some sites other than the nares, such as the lungs, as being the entry point of S1 detected in blood.

It is important to note that although the study shows that I-S1 crosses the BBB in mice, this may not be the case in humans. For that reason, we used in vitro models of the human BBB, which can be useful in studying mechanisms of BBB permeability. The model used in this study is derived from human iPSCs and develops a brain endothelial cell-like phenotype that includes functional BBB influx and efflux transporters and strong barrier properties that permit the study of transport without confounding effects of high baseline leakage.

In this model, we did not observe significant differences in permeability for I-S1 compared to T-Alb. The apparent absence of I-S1 transport across the BBB in this in vitro model could be due to technical issues, such as blockers of I-S1 binding in the buffers. It could also mean that the iBECs did not express the cell-membrane glycoproteins necessary for I-S1 transport, or that I-S1 is not able to cross the human BBB. A note of caution regarding the validity of using monomeric S1 as a model for SARS-CoV-2 is that S1 is normally attached to SARS-CoV-2 as a trimer. However, the S1 protein may be shed from the virus in vivo, and therefore studying S1 monomers may...
have validity by itself—although there is currently no direct evidence that spike proteins are shed from SARS-CoV-2. Altogether, our results strongly suggest that the S1 protein can cross the murine BBB through a mechanism resembling adsorptive transcytosis and be taken up by peripheral tissues independently of human ACE2.

Online content
Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41593-020-00771-8.

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Methods

Mice. All mouse studies were approved by the local Institutional Animal Care and Use Committee at Oregon Health & Sciences University and the Veterans Affairs Puget Sound Health Care System (VAPSHCS) and conducted in facilities approved by the American Association for Accreditation of Laboratory Animal Care (AAALAC). Male CD-1 mice (6–10 weeks old) were purchased from Charles River Laboratories (Hollister). A total of 204 CD-1 males were used for all studies. Human E3- and E4-targeted replacement (TR) male and female mice, generated as described previously38, were bred at the Oregon Health & Sciences University before transfer to the VAPSHCS for the experiments. E3 and E4 TR mice were approximately 4 months of age on the day of the study. A total of 21 E3 and 20 E4 TR mice were used for all studies. Mice had ad libitum access to food and water and were kept on a 12:12 h light/dark cycle. The temperature was maintained at 18–23°C and humidity was maintained at 40–60%. For all experiments, mice were anesthetized with an intraperitoneal injection of 0.15–0.2 ml of 40% urethane to minimize pain and distress. Anesthetized mice were placed on a heating pad until time of use. At the end of each study, mice were euthanized by decapitation while under anesthesia.

Sources and radioactive labeling of proteins. The S1 proteins were provided by the manufacturers (RayBiotech, 230-30161, Val 16–Glino 690, AMSBO, AMS, SIN-S253H, Val 16–Arg 685). The proteins were dissolved in PBS (pH 7.4) at a concentration of 0.45–0.61 mg ml⁻¹. The proteins were produced in HEK293 cell lines, had C-terminal His tags, had C-terminal His tags, and either were immediately or stored at –80 °C until use. The 5 μg of thawed S1 protein was radioactive labeled with 1 μCi 111In (Perkin Elmer) using the chloramine-T method, as described previously38. The I-S1 was purified on a column of G-10 Sephadex (GE Healthcare) and eluted with PBS into glass tubes containing 1% BSA in lactated Ringer’s solution (BSA-LR). We estimated based on the amount of protein iodinated that the specific activity of I-S1 was approximately 11 Ci g⁻¹ or about 12.5 ng per 300,000 c.p.m. BSA (Sigma) was labeled with 99mTc (GE Healthcare) using the stannous tartrate method36. T-Alb was purified on a column of G-10 Sephadex. Both of the I-S1 and the T-Alb were more than 90% acid precipitable. The molecular weight of the labeled proteins was confirmed by running 200,000-600,000 c.p.m. activity in 1× LDS buffer (Invitrogen) with or without reducing agent (Invitrogen) on a 4–12% Bis-Tris gel (GenScript) in MOPS buffer (Invitrogen). The gel was then fixed for 30 min in 10% acetic acid/50% methanol, washed 3× with water and then dried using a DryEase Mini-Gel Drying System (Invitrogen). Dried gels were exposed on autoradiography film for 24 h and then developed. Major bands of I-S1 from RayBiotech and AMSBO migrated at their predicted molecular weight patterns, based on the manufacturer’s data (Extended Data Fig. 1).

Measurement of blood-to-brain entry rate. MRTR34,35 was used to measure the unidirectional blood-to-brain influx constant (K) for I-S1. In anesthetized mice, the left jugular vein was exposed for an i.v. injection of 0.1 ml BSA-LR containing 3 × 10⁵ c.p.m. of I-S1. T-Alb (6 × 10⁵ c.p.m.) was also included in the injection for measurement of the vascular space for the brain and the albumin space for the liver. The percentage of radioactivity in blood from 0–30 min was calculated using Prism software (GraphPad). Results for serum were expressed as the percentage of the injected dose per ml of blood (% inj ml⁻¹). The percentage of the injected dose per gram of brain (% inj g⁻¹) was calculated by multiplying the delta brain/serum value by the percentage of injected dose per ml for I-S1.

Stability of I-S1 in brain and blood. Anesthetized mice received an injection into the jugular vein of 0.1 ml of BSA-LR containing 6 × 10⁵ c.p.m. of I-S1 and 6 × 10⁵ c.p.m. of T-Alb. Ten minutes after the injection, arterial blood was collected from the abdominal aorta, the thorax opened and the descending thoracic artery clamped, both jugular veins severed and 20 ml of lactated Ringer’s solution perfused through the left ventricle of the heart to wash out the vascular space of the brain. The whole brain was then removed. Whole blood was centrifuged and 10 μl of serum was added to 500 μl of BSA-LR and combined with an equal part of 30% trichloroacetic acid, mixed, centrifuged and the supernatant and pellet counted. Whole brains were homogenized in BSA-LR with complete mini protease inhibitor cocktail (Roche) and then homogenized with a handheld glass homogenizer and centrifuged. The resulting supernatant was combined with equal parts of 30% trichloroacetic acid, mixed, centrifuged and the supernatant and pellet counted. To determine the percentage of degradation of I-S1 or T-Alb that occurred during processing, I-S1 and T-Alb were added to brains and arterial whole blood from animals that had not been injected with radioactivity and processed immediately as above. The percentage of radioactivity that was precipitated by acid (%P) in all of these samples was calculated by the equation:

\[
\%P = 100\left(\frac{S}{S + P}\right)
\]

where S is the c.p.m. in the supernatant and P is the c.p.m. in the pellet.

Capillary depletion of brain. The capillary depletion method as adapted to this assay used to separate cerebral capillaries and vascular components from brain parenchyma53,54. We used the variant of the technique that also estimates reversible binding to the capillary lumen. The latter is done by assessing two groups of mice: one with a vascular washout step, which removes material reversibly bound to the capillary lumen (washout group), and one without the washout step (non-washout group). The difference between these groups represents material that was reversibly adhering to the capillary lumen (equation 3). Mice were anesthetized and received an i.v. injection of 6 × 10⁵ c.p.m. T-Alb with 6 × 10⁵ c.p.m. I-S1 in 0.1 ml BSA-LR. At 5, 10 and 30 min later, blood was obtained from the carotid artery, and the brain (non-washout group) was extracted. In a separate group of mice, blood was taken from the abdominal aorta at 5, 10 and 30 min, the thorax was opened and the descending thoracic artery clamped, both jugular veins severed and 20 ml of lactated Ringer’s solution infused via the left ventricle of the heart to wash out the vascular contents of the brain and any material reversibly associated with the capillary lumen (washout group). Each whole brain was homogenized in glass with physiological buffer (10 mM HEPES, 141 mM NaCl, 4 mM KCl, 2.8 mM CaCl₂, 1 mM MgSO₄, 1 mM NaH₂PO₄, H₂O, 10 mM d-glucose (pH 7.4)) and mixed thoroughly with 26% dextran. The homogenate was centrifuged at 4,255 × g for 15 min at 4°C. The pellet, containing the capillaries, and the supernatant, representing the brain parenchymal space, were carefully separated. Radioactivity levels in the capillary pellet, the brain supernatant and the arterial serum were determined for both T-Alb and I-S1 and expressed as the capillary/serum and brain parenchyma/serum ratios. The I-S1 parenchymal ratios were corrected for vascular contamination by subtracting the corresponding ratios for T-Alb; these results are reported as the delta brain parenchyma/serum ratios. The amount of S1 in the brain parenchymal space was taken as the delta brain parenchymal space from the washout group (P_b), the amount in the capillary as the capillary from the washout group (C_c), and the amount of material loosely binding to the luminal surface (luminal) as:

\[
\text{Luminal} = (P + C) - (P_b + C_c)
\]

Where P is the delta brain parenchymal space and C is the capillary space, both from the non-washout groups.

Effects of wheatgerm agglutinin and heparin. We co-injected WGA with I-S1 to determine the role of glycoproteins containing stalic acid and N-acetylgalosamine in the uptake of S1 by the BBB and other tissues. Mice were anesthetized, after which the jugular vein and right side of the heart were exposed. Mice then received a jugular venous injection of 0.1 ml of BSA-LR containing 3 × 10⁵ c.p.m. of I-S1 and 6 × 10⁵ c.p.m. of T-Alb and, in a subset of mice, 10 μg of the plant lectin WGA (Sigma). Brain, tissues and serum samples were collected 5 min later and tissue/serum ratios calculated as above in units of μl per gram. Heparin sulfate is used as a receptor by some viruses19 and heparin by binding to viral proteins can block viral entry into brain19. Therefore, separate groups of mice had heparin (12 U per mouse) included in the injection.

Saturation of I-S1 and binding to ACE2 enzyme. In anesthetized mice, the left jugular vein i.v. injection of 0.1 ml of BSA-LR containing 3 × 10⁵ c.p.m. of I-S1 (RayBiotech) and 6 × 10⁵ c.p.m. of T-Alb. For some mice, the injection contained 1 μg per mouse of unlabeled S1 (AMSBO), mouse aeryl ghrilin (CBo), angiotensin II (ToCris) or human ACE2 (R&D). Ten minutes after i.v.
injection, whole blood was obtained from the carotid artery and centrifuged after clotting. The whole brain, olfactory bulb, kidney, spleen and portions of the liver and lung were removed. The levels of radioactivity in the arterial serum and the tissues were determined and the results expressed as the percentage of the injected I-S1 per ml for serum and delta tissue/serum ratios for the tissues.

Lipopolysaccharide injections. Male CD-1 mice aged 6–10 weeks were given an intraperitoneal injection of 5 mg kg \(^{-1}\) LPS from S. typhimurium (Sigma) dissolved in sterile saline at 0, 6, and 24h. At 28h, mice were anesthetized with an intraperitoneal injection of 60 mg kg \(^{-1}\) Xylazine (Prior, 1001) and 1 mg kg \(^{-1}\) Ketamine (HESFM; Sigma, 11111044) supplemented with 20 ng ml \(^{-1}\) bFGF, 10 µM retinoic acid and 1% B27. At 24h after subcutaneous injection of bFGF, 10 µM retinoic acid and 1% B27. At 24h after subcutaneous injection, the medium was changed to HESFM + 1% B27 without bFGF or retinoic acid, and transendothelial electrical resistance (TEER) was recorded using an End EVO20 Voltohmmeter (World Precision Instruments) coupled to an EndOhm cup chamber. TEER measurements occurred daily and S1 transport experiments were conducted when TEER stabilized, between 10–13 d in vitro. TEER values in these studies exceeded 1,000 Ω cm \(^{-2}\), and TEER mean values were confirmed to be equal among groups just before starting the transport study. Before transport studies, the medium was changed and cells were equilibrated in the incubator for 20 min. Warm HESFM + 1% B27, 1 million c.p.m. of T-Alb and 500,000 c.p.m. of I-S1 were then added in a volume of 100 µl to the luminal chamber. After incubation times of 10, 20, 30 and 45 min at 37 °C, 500-µl volumes of medium from the abluminal chamber were collected and replaced with fresh prewarmed medium. Samples were then acid precipitated by adding a final concentration of 1% BSA to visualize the pellet and 15% trichloroacetic acid to precipitate the proteins in solution. The samples were centrifuged at 4,255g for 15 min at 4 °C. Radioactivity in the pellet was counted in the gamma counter and the permeability surface-area coefficients for T-Alb and I-S1 were calculated according to the method by Dehouck et al. \(^{54}\). Clearance was expressed as µl of radioactive tracer that was transported from the luminal to abluminal chamber, calculated from the initial level of acid-precipitable radioactivity added to the luminal chamber and the final level of radioactivity in the abluminal chamber:

\[
\text{Clearance(µl) = } \frac{C_l + VC_s}{C_i} \times \frac{1}{PS_{app}}
\]

Where \(C_l\) is the initial concentration of radioactivity in the luminal chamber (in c.p.m. µl \(^{-1}\)), \(C_s\) is the concentration of radioactivity in the abluminal chamber (in c.p.m. µl \(^{-1}\)), \(V_s\) is the volume of the serum and \(V_c\) is the volume of the transwell insert. Clearance was divided by the surface area of the Transwell inserts (0.33 cm \(^2\)) to generate the endothelial permeability coefficient (\(P_{f}^{e}\) in µl min \(^{-1}\) cm \(^{-2}\)).

Statistical and reproducibility. Sample size, randomization and blinding. Sample sizes were predetermined based on the variance that is known from previously conducting similar experiments on BBB transport. We selected samples sizes to provide a power of 0.8–0.9 based on 30–40% differences between means. Anesthetized mice were randomized between test groups by alternating their assignments. Transwells were assessed for TEER before experiments and assigned to groups in such a way that TEER values between groups had approximately equal means and variances. Blinding was not done for experiments or analyses because this knowledge was required to carry out experiments and analysis.

Data exclusion and replication. For MTRA, the method calls for exclusion of data points that contribute to nonlinearity of the curve. Outliers whose exclusion improved the \(r^2 ≥ 0.2\) were excluded from the analysis (excluded points are denoted by filled circles in Figs. 2 and 3 and further explained in the legends). For ANOVAs and comparisons of means, the Grubbs outlier exclusion test (alpha < 0.05) was applied once to detect single outliers, and any outliers that were excluded by this method were stated in the figure legends and their values provided. Each figure was generated as a single experimental replicate, with the exception of iPSC studies where S1 and albumin transport were compared in three independent experiments. However, use of T-Alb as an internal control for all experiments allowed us to track consistency between experiments with different designs based on its predictable tissue/serum ratios. Further, the I-S1 tissue/serum ratios taken at the same time points from different experiments could be compared, and agreed well with each other, demonstrating a reproducible observation of I-S1 transport into the brain and other tissues.

Statistical analysis. Prism 8.0 was used for all statistical calculations (GraphPad). Simple linear regression analysis was used to calculate slopes and intercepts, and the error terms for the slope and intercept were the standard errors. As required by MTRA, only the linear portion of the slope was used to calculate \(K_c\). ANCOVA was used to determine whether slopes and intercepts of two lines differed, and two-way ANCOVA was used to determine effects of sex and genotype on regression lines in the APOE mouse studies. Two-tailed t-tests were used to compare two means, and a one-way or two-way ANOVA and multiple-comparisons testing were used when more than two means were compared. The statistical tests used are specified in all figure legends. Data distribution was assumed to be normal, but this was not formally tested. However, all data points are shown with the mean and standard deviation.

Reporting Summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability. The data that support the findings of this study are available from the corresponding author upon reasonable request. Source data are provided with this paper.
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Author contributions

E.M.R.: experimental design, experimental work and data analysis (mouse experiments), writing, editing and graphics. A.F.L.: writing, editing, experimental work and data analysis (mouse experiments). K.M.H.: writing, editing, experimental work and data analysis (protein labeling and characterization and mouse experiments). L.M.W.: experimental work and data analysis (in vitro studies), writing and editing. M.J.R.: experimental work (mouse experiments), experimental design, writing and editing. K.K.B.: experimental work (APOE mouse husbandry), writing and editing. J.R.: writing, editing and experimental design. W.A.B.: experimental design, statistics, writing, editing and graphics. M.A.E.: writing, editing, experimental design and experimental work (protein labeling, mouse experiments and in vitro studies).

Competing interests

The authors declare no competing interests.

Additional information

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Extended Data Fig. 1 | Gel Autoradiography of I-S1 from RayBiotech and AMSBIO. Cropped images of Fraction 9 of I-S1 from RayBiotech (left panel, 1μl of approximately 300,000 CPM run under reducing and non-reducing conditions, exposed to film for 24 hours), and from AMSBIO (right panel, 1μl of approximately 600,000 CPM run under reducing and non-reducing conditions. In each case, both the reduced and non-reduced gels migrated at the molecular weights predicted by the manufacturers. The autoradiography gels to validate size were run once. The quality of subsequent I-S1 labeling was confirmed by acid precipitation, described in methods.
Extended Data Fig. 2 | I-S1 is uniformly taken up by all brain regions. Mice were given an injection of I-S1 (RayBiotech) and olfactory bulb, whole brain, and blood were collected 10 min later. Brains were dissected into the 10 regions shown. One-way ANOVA with brain regions as the independent variable showed a trend for differences (p = 0.09) and post-hoc multiple comparisons tests showed no differences among the brain regions. N = 11 mice. FCtx = frontal cortex, PCtx = parietal cortex, OCtx = occipital cortex.
Extended Data Fig. 3 | Capillary depletion for I-S1. Capillary depletion studies were conducted to determine partitioning of the I-S1 (RayBiotech) in the capillary lumen, in the capillary, and in the brain parenchyma over time. The compartment that contained radioactivity was plotted against clock time, and the slopes were analyzed by linear regression. The slope of the line plotted for the parenchymal fraction was $0.187 \pm 0.016 \, \mu L/g \cdot min$, $r^2 = 0.952$, $n = 9$ mice, and was non-zero ($p < 0.0001$); the slope for the capillary fraction was $0.027 \pm 0.006 \, \mu L/g \cdot min$, $r^2 = 0.757$, $n = 9$ mice, and was non-zero ($p = 0.0023$). The slope for luminal binding was $0.05041 \pm 0.04555 \, \mu L/g \cdot min$ and did not significantly deviate from zero ($p = 0.3050$, $n = 9$, $r^2 = 0.145$). The mean value for luminal binding over all time points was $1.75 \pm 0.50 \, \mu L/g$. 

![Graph showing partitioning of I-S1 in different compartments over time](image-url)
Extended Data Fig. 4 | Transport of S1 proteins across human iPSC-derived brain-like endothelial cell (iBEC) monolayers. a. RayBiotech I-S1 passage across the iBEC monolayer was not significantly faster than that of T-Alb (p = 0.1464, t = 1.527, df=16; two-tailed paired t-test). Combination of 3 independent differentiations, n = 17 transwells/group, average TEER (2578 ± 897.8 Ω·cm²). b. Transport of RayBiotech I-S1 in the presence of 5 µg/ml excess unlabeled S1. Excess S1 had no significant effect on I-S1 or T-Albumin Pe (p > 0.5; two-way ANOVA). N = 5-6 transwells/group from a single differentiation. c. Transwells were treated with WGA (0.5 mg/ml) or vehicle. There was a significant overall effect of treatment (F(1, 9) = 9.435, p = 0.0133) and analyte (F(1, 9) = 6.949, p = 0.0271) on RayBiotech I-S1 or T-Alb transport (two-way ANOVA), but there were no significant effects of WGA on I-S1 or T-Alb Pe (Sidak’s multiple comparisons testing). N = 5-6 wells/group from a single differentiation. d. Comparison of RayBiotech I-S1 and AMSBIO I-S1 transport. T-Alb was included with each I-S1 and also compared to the I-S1. Two-way ANOVA showed differences between I-S1s from the two sources (F(1,10) = 10.1, p = 0.0098), but not between analytes (T-Alb vs I-S1). Multiple comparisons using Sidak’s test showed a faster transport for RayBiotech I-S1 vs AMSBIO I-S1 (p = 0.0324). N = 6 wells/group from a single differentiation. *p < 0.05.
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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

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| Data collection | No software used |
|-----------------|------------------|
| Data analysis   | Prism 8.0 (GraphPad, Inc, San Diego CA) was used for data analysis |

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | Variance is known from previously conducting similar experiments and power at 0.8-0.9 based on 30-40% differences between means |
|-------------|--------------------------------------------------------------------------------------------------------------------------|
| Data exclusions | For multiple-time regression analysis, the method calls for exclusion of data points that contribute to non-linearity of the curve. Outliers whose exclusion improved the $r^2 > 0.2$ were excluded from analysis. Excluded points are noted in Figures 2 and 3 by filled circles and further explained in those figure legends. For ANOVAs and comparisons of means, the Grubbs outlier exclusion test (alpha < 0.05) was applied once to detect single outliers, and any outliers that were excluded by this method are stated in the Figure legends and their values provided. |
| Replication | Each figure was generated as a single experimental replicate, with the exception of iPSC studies where S1 and albumin transport were compared in three independent experiments. However, use of T-Alb as an internal control for all experiments allowed us to track consistency between experiments with different designs based on its predictable tissue/serum ratios. Further, the I-S1 tissue-serum ratios taken at the same time points from different experiments may be compared, and agree well with each other, demonstrating a reproducible observation of I-S1 transport into the brain and other tissues. |
| Randomization | Anesthetized mice were randomized between test groups by alternating their assignments. Transwells were assessed for TEER prior to experiments and assigned to groups in such a way that TEER values between groups had approximately equal means and variances. |
| Blinding | Blinding was not done for experiments or analysis because this knowledge is required to carry out experiments and analysis. |

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We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

| Materials & experimental systems | Methods |
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| Antibodies | Involved in the study |
| Eukaryotic cell lines | ChIP-seq |
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| Clinical data | |
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Eukaryotic cell lines

Policy information about cell lines

- Cell line source(s): GM25256 iPSC line from the Coriell Institute
- Authentication: Cells are routinely verified for iPSC markers such as Oct4, Nanog, and alkaline phosphatase
- Mycoplasma contamination: Cell lines were not tested for mycoplasma contamination in-house
- Commonly misidentified lines (See ICLAC register): Not applicable

Animals and other organisms

Policy information about studies involving animals; ARRIVE guidelines recommended for reporting animal research

- Laboratory animals: 6-10 week-old male CD-1 mice, and 4 month old male and female APOE3 and APOE4 targeted replacement mice.
| Wild animals          | Wild animals were not used          |
|----------------------|------------------------------------|
| Field-collected samples | Field-collected samples were not used. |
| Ethics oversight     | Studies were approved by the institutional animal research review board and the facilities are approved by AAALAC |

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