Endothelial TLR4 and the microbiome drive cerebral cavernous malformations

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Cerebral cavernous malformations (CCMs) are a cause of stroke and seizure for which no effective medical therapies yet exist. CCMs arise from the loss of an adaptor complex that negatively regulates MEKK3–KLF2/4 signalling in brain endothelial cells, but upstream activators of this disease pathway have yet to be identified. Here we identify endothelial Toll–like receptor 4 (TLR4) and the gut microbiome as critical stimulants of CCM formation. Activation of TLR4 by Gram–negative bacteria or lipopolysaccharide accelerates CCM formation, and genetic or pharmacologic blockade of TLR4 signalling prevents CCM formation in mice. Polymorphisms that increase expression of the TLR4 gene or the gene encoding its co–receptor CD14 are associated with higher CCM lesion burden in humans. Germ–free mice are protected from CCM formation, and a single course of antibiotics permanently alters CCM susceptibility in mice. These studies identify unexpected roles for the microbiome and innate immune signalling in the pathogenesis of a cerebrovascular disease, as well as strategies for its treatment.

CCMs are relatively common vascular malformations that arise predominantly in the central nervous system, causing haemorrhagic stroke and seizure1. CCMs arise from loss of function mutations in three genes, KRIT1, CCM2 and PDCD10, that encode components of a heterotrimeric, intracellular adaptor protein complex (the ‘CCM complex’)2–3. The clinical course of familial CCM disease is highly variable, even among individuals who share identical germline mutations4–6, suggesting the existence of powerful genetic and/or environmental disease modifiers. Present treatment for CCMs consists solely of palliative therapies or neurosurgical resection.

Previous studies of vertebrate genetic models and human CCM lesions have demonstrated that loss of the CCM complex results in vascular lesion formation owing to increased MEKK3–KLF2/4 signalling in brain endothelial cells7–9, and that the CCM complex suppresses MEKK3–KLF2/4 signalling through a direct interaction between CCM2 and MEKK3 (refs 10, 11). As there is a lack of effective drugs that target the MEKK3–KLF2/4 pathway, these molecular insights have not been immediately translational. However, they raise a key mechanistic question: if the role of the CCM complex is to negatively regulate MEKK3–KLF2/4 signalling, what activates this pathway in brain endothelial cells? Identification of upstream activators of this pathway is needed to understand the pathogenesis of CCM disease and reveal viable therapeutic strategies.

**CCM formation is driven by GNB and LPS**

To investigate CCM formation in mice, we generated animals in which endothelial-specific deletion of Kr1t1 or Ccm2 was induced one day after birth (P1, iEcrt1Δ/Δ and iEcrt1ΔCcm2Δ/Δ, hereafter denoted as Kr1t1ECKO and Ccm2ECKO, respectively). In this model, vascular malformations first appear in the cerebellar white matter at P6, with numerous mature lesions present by P10 (refs 9, 13). These mice were maintained as inbred breeding colonies and initially demonstrated a highly penetrant lesion phenotype (termed ‘susceptible’; Fig. 1a, top). However, following a change in vivarium at the University of Pennsylvania, we noted the spontaneous emergence of Kr1t1ECKO and Ccm2ECKO sub–colonies that developed barely visible hindbrain lesions at P17 (termed ‘resistant’; Fig. 1a, bottom). Previous studies have demonstrated 100% CCM penetrance on a C57BL/6J background13, but Ccm2ECKO animals backcrossed seven generations to C57BL/6J remained resistant to CCM formation (Extended Data Fig. 1a). Importantly, among a large population of CCM-resistant animals, we detected a small number of individual pups that exhibited robust CCM formation in association with the presence of intra-abdominal, Gram-negative bacterial (GNB) abscesses that probably developed following tamoxifen injection (Fig. 1b, c, Extended Data Fig. 1b). This observation suggested that Gram-negative infection accelerates CCM pathogenesis.

To directly assess the role of Gram-negative infection, Gram-negative abscesses were induced in resistant Ccm2ECKO mice at P5 by intra-peritoneal injection of live *Bacteroides fragilis*. Following *B. fragilis* injection, 9 out of 16 resistant Ccm2ECKO animals developed large CCM lesions (termed ‘responders’; Fig. 1d, left) and 7 out of 16 animals did not (termed ‘non-responders’; Fig. 1d, right). Responders to *B. fragilis* injection exhibited splenic abscesses and higher spleen weights compared with non-responders (Fig. 1e, f), suggesting that
Susceptible through the innate immune receptor TLR4 (ref. 14), and MEKK3- signalling in CCM-complex-deficient brain endothelial cells. Analysis of LPS accelerate CCM formation by activating TLR4–MEKK3–KLF2/4 responses to LPS in vitro signalling in endothelial cells 9. LPS activates intracellular signals findings reveal that blood-borne GNB and LPS are strong drivers of formation of large lesions in resistant of LPS to drive CCM formation. Injection of LPS resulted in the formation in resistant Ccm2 required to stimulate CCM formation in resistant unpaired, two-tailed ANOVA with Holm–Sidak correction for multiple comparisons or unpaired, two-tailed t-test. ****P < 0.0001; ***P < 0.001; **P < 0.01; NS, P > 0.05.

**Figure 1** | CCM formation is stimulated by GNB infection and intravenous LPS injection. a, Lesion formation in susceptible and resistant Ccm2ECKO mice at P17. Dotted lines trace cerebellar white matter. Asterisks, CCM lesions; scale bars, 1 mm. b, Hindbrains of resistant Ccm2ECKO littermates without (top) and with (bottom) spontaneous abdominal Gram-negative abscess. Arrows, CCM lesions; scale bars, 1 mm. c, The bacterial abscess (‘absc’) identified in b contains GNB (arrows). Scale bars, 4 mm (top) and 10 μm (bottom). d, CCM formation in resistant Ccm2ECKO littermates following infection with a live B. fragilis/autochlorae caecal contents mixture (B. fragilis) or autoclaved caecal contents (ACC) alone (ACC vehicle). Scale bars, 1 mm. e, f, Resistant Ccm2ECKO responders exhibit splenic abscesses and increased spleen weight compared with non-responders. g, CCM formation in resistant Ccm2ECKO mice following vehicle or LPS treatment. Scale bars, 1 mm. h, i, Quantification of lesion and total brain volumes. Error bars shown as s.e.m. and significance determined by one-way ANOVA with Holm–Sidak correction for multiple comparisons. ****P < 0.0001; ***P < 0.001; **P < 0.01; NS, P > 0.05.

Endothelial TLR4 drives CCM lesion formation

We previously demonstrated that loss of the CCM proteins Krit1 or CCM2 results in vascular malformation owing to increased MEKK3 signalling in endothelial cells. LPSactivates intracellular signals through the innate immune receptor TLR4 (ref. 14), and MEKK3-deficient fibroblasts are unable to activate downstream signalling responses to LPS in vitro. We therefore hypothesized that GNB and LPS accelerate CCM formation by activating TLR4–MEKK3–KLF2/4 signalling in CCM-complex-deficient brain endothelial cells. Analysis of susceptible Krit1ECKO mice revealed that CCM lesions arise in the absence of an immune-cell infiltrate at P6 (Extended Data Fig. 2), and that a single dose of LPS at P5 accelerated CCM formation by P6 (Fig. 2a). Consistent with a mechanism that is intrinsic to brain endothelial cells, we observed synergistic effects of CCM-complex-deficiency and LPS injection on the expression of CCM-driving genes Klj2 and Klj4, known endothelial TLR4 signalling targets IL-1β (IL1β) and E-selectin (Sele), as well as on the level of phospho-myosin light chain (Fig. 2b, Extended Data Fig. 1e).

To directly assess the requirement for endothelial TLR4 in spontaneous CCM formation, we bred iECre;Krit1flo/flo;Tlr4flo/flo and iECre;Krit1flo/flo, Tlr4flo/flo mice using animals from the susceptible Krit1ECKO colony. Loss of a single endothelial Tlr4 allele resulted in an approximately 75% reduction in CCM lesion burden at P10, whereas loss of both resulted in virtually complete prevention of CCM lesion formation (Fig. 2c, d, Extended Data Fig. 3a). Cd14 encodes a soluble TLR4 co-receptor that binds LPS and facilitates TLR4 signalling. Although less complete, global loss of Cd14 also prevented CCM formation in susceptible Krit1ECKO mice (Fig. 2e, f, Extended Data Fig. 3b). Lineage tracing studies confirmed that Cdh5(PAC)-CreERT2 transgene activity was restricted to endothelial cells (Extended Data Fig. 4), excluding a requirement for haematopoietic cell TLR4 signalling during CCM formation. Finally, to exclude a role for the CCM complex in endothelial cells outside of the brain, we used a recently generated SloCl1(BAC)-CreERT2 transgene to further restrict deletion of Krit1 to

**Figure 2** | CCM lesion formation requires endothelial TLR4/CD14 signalling. a, Injection of LPS at P5 drives CCM formation by P6 in susceptible Krit1 ECKO littermates. Scale bars, 1 mm (white) and 50 μm (yellow); arrows and arrowheads, CCM lesions; dotted lines, cerebellar white matter. b, Gene expression in cerebellar endothelial cells isolated from the indicated littermates at P6. n ≥ 3 per group. c–f, Genetic rescue of CCM formation with endothelial loss of TLR4 or global loss of CD14. Visual appearance of CCM lesions (above), corresponding X-ray micro-computed tomography (microCT) images (below), and lesion volume quantification. Scale bars, 1 mm. Error bars shown as s.e.m. and significance determined by one-way ANOVA with Holm–Sidak correction for multiple comparisons. ****P < 0.0001; ***P < 0.001; **P < 0.01; NS, P > 0.05.

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Figure 3  |  Increased TLR4 or CD14 expression is associated with higher lesion number in familial CCM patients. a, SNPs in the 5’ genomic regions of TLR4 and CD14 associated with increased lesion numbers in patients with familial CCM are shown relative to the transcriptional start site (TSS). b, c, Normalized microarray measurement of TLR4 and CD14 expression in whole-blood cells from individuals in the general population and with the indicated CD14 expression in whole-blood cells from individuals in the general population with the indicated TLR4 and CD14 genotypes, respectively. d, Representative MRI images of KRIT1(Q455X)-carrying patients with raw lesion count and TLR4 and/or CD14 SNP genotypes (RA, risk allele). Arrow indicates a CCM lesion. e, f, Sex- and age-adjusted log(lesion burden) in KRIT1(Q455X)-carrying patients with indicated the genotypes. Error bars shown as 95% confidence intervals and significance determined by one-way ANOVA with Holm–Sidak correction for multiple comparisons. ***P < 0.0001; **P < 0.001; *P < 0.01; *P < 0.05.

brain endothelial cells. Scl01c(BAC)-CreER(T2)-R26-LSL-RFP animals, in which cellular Cre activity is marked by expression of red fluorescent protein (RFP), exhibited RFP+ endothelial cells in the brain but not in the gut or liver (Extended Data Fig. 5a), and Scl01c(BAC)-CreER(T2), Krit1(ECKO) animals developed CCM lesions like those in Krit1(ECKO) animals (Extended Data Fig. 5b, c). These genetic findings identify endothelial TLR4 signalling as a critical driver of CCM formation in mice.

**TLR4/CD14 expression parallels human CCM burden**

Studies in humans and mice have demonstrated that TLR4 signalling positively correlates with receptor expression levels, suggesting that polymorphisms associated with changes in TLR4 expression might influence the progression of human CCM disease. We recently analysed 830 genetic variants of 56 inflammatory and immune related genes in 188 patients with an identical nonsense mutation in the KRIT1 gene (Q455X) in whom CCM lesion burden was measured using magnetic resonance imaging (MRI). Following statistical analysis, single-nucleotide polymorphisms (SNPs) in only two genes, TLR4 (rs10759930, chromosome 9; Fig. 3a) and CD14 (rs7785878, chromosome 5; Fig. 3a), were found to be significantly associated with increased CCM lesion number. Further analysis of genes in TLR4–MEKK3–KLF2/4 signalling pathways identified additional SNPs for TLR4 (rs10759931) and CD14 (rs778588) in linkage disequilibrium with those previously identified (Fig. 3a), but none in other pathway genes (See Methods) that associated with altered lesion burden.

Notably, the TLR4 and CD14 SNPs associated with increased CCM lesion number are in the 5’ genomic region of each gene (Fig. 3a), and constitute cis expression quantitative trait loci (cis-eQTLs) that positively regulate whole-blood-cell expression of TLR4 and CD14 in a dose-dependent manner corresponding with risk allele number.

**Bacteria drive CCM formation in mice**

Although endogenous TLR4 ligands have been identified, the best-characterized TLR4 ligand is GNB-derived LPS. The findings that CCM pathogenesis requires endothelial TLR4 and CD14 (Fig. 2c–f), and that CCM susceptibility shifted markedly with a change in vivarium (Fig. 1a), suggested that GNB in the microbiome may be a primary source of TLR4 ligand and an important regulator of CCM disease. To directly assess the role of the bacterial microbiome during CCM formation, we delivered susceptible E19.5 Krit1(ECKO) neonates using sterile caesarean section (C-section) and fostered them to imported conventional or germ-free Swiss Webster mothers (Fig. 4a). All fostered Krit1(ECKO) neonates exhibited robust CCM formation at P10 when raised by conventional Swiss Webster mothers (Fig. 4b, c, Extended Data Fig. 3c). In contrast, 7 out of 8 fostered Krit1(ECKO) neonates raised in germ-free conditions failed to develop CCM lesions, indicating that bacteria are required for CCM pathogenesis in most animals (Fig. 4b, c, Extended Data Fig. 3c). A single fostered Krit1(ECKO) neonate developed CCM lesions despite reductions in gut bacteria and Krit1 mRNA, similar to fostered Krit1(ECKO) littermates that failed to develop CCMs (red boxes, Fig. 4b–e, Extended Data Fig. 3c). Previous studies have demonstrated that MEK3 is required for signalling downstream of cytokines IL-1β and TNFα, and other pattern-recognition receptors can signal in endothelial cells through the same effectors used by TLR4 (refs 26, 27). Thus, the generation of lesions in a germ-free Krit1(ECKO) neonate suggested that cytokines or innate receptors other than TLR4 may also drive CCM formation in vivo. To directly test the ability of non-TLR4 ligands to stimulate CCM formation, we administered IL-1β and TNFα to ECKO neonates using sterile caesarean section (C-section) and fostered them to imported conventional or germ-free Swiss Webster mothers (Fig. 4a). Lesion numbers in ECKO neonates and fostered them to imported conventional or germ-free Swiss Webster mothers significantly increased CCM lesion volume, although no difference was observed with TNFα or peptidoglycan (Extended Data Fig. 6). These findings identify the bacterial biome as a critical driver of CCM formation in vivo, but also demonstrate that cytokines and innate immune ligands other than LPS can support CCM formation in vivo.

**CCM susceptibility is associated with gut GNB**

Assessment of CCM formation in Krit1(ECKO) and Ccm2(ECKO) mice at P10 revealed a remarkably binary phenotype in which susceptible mice developed numerous lesions, whereas resistant mice developed no lesions. To identify specific bacteria that associate with CCM susceptibility or resistance, we performed 16S rRNA gene sequencing of bacterial
DNA extracted from the faeces of female mice that raised susceptible or resistant Krit1ECKO and Ccm2ECKO animals (Extended Data Fig. 7a).

A PERMANOVA test of unweighted UniFrac distances (a measure of difference between biological groups) revealed clear separation of susceptible and resistant bacterial microbiome communities, regardless of whether they were derived from Krit1ECKO or Ccm2ECKO colonies ($P < 0.0001$, $R^2 = 0.051$; Fig. 5f). Further accounting for relative abundances of bacterial species, significant separation between susceptible and resistant animals was also observed using weighted UniFrac analysis ($P = 0.0016$, $R^2 = 0.091$; Fig. 5g). Fitting generalized, linear mixed-effects models for commonly present bacterial taxa identified one major group that differed significantly between the gut biomes of susceptible and resistant animals: Gram-negative Bacteroidetes family s24-7 (denoted as s24-7) was significantly more abundant in susceptible animals irrespective of genotype (Fig. 5h, Extended Data Fig. 7b, c). Notably, 16S sequencing of gut bacteria from conventional Swiss Webster foster mothers revealed high levels of s24-7, explaining the lower CCM lesion volume in conventional Swiss Webster (SW) foster mothers compared to Krit1ECKO and Ccm2ECKO mothers ($P < 0.0001$; Fig. 5b). These findings support a key role for the gut microbiome 28–31. This model predicts two novel approaches to treat CCM disease: TLR4 blockade and manipulation of the gut microbiome.

**TLR4 block or altering the microbiome prevents CCM**

Our studies do not exclude a role for TLR4 signalling in non-brain endothelial cells. However, they are most consistent with a disease model in which brain endothelial TLR4 and/or CD14 receptors interact with s24-7 to promote microvascular angiogenesis. Krit1ECKO or Ccm2ECKO animals fostered to conventional Swiss Webster (SW) mothers. Significance determined by linear mixed effects model with Benjamini–Hochberg correction for multiple comparisons. $**P < 0.0001$; $***P < 0.001$; $****P < 0.05$. Note, conventional Swiss Webster data from Figs 4 and 5 are the same experiment.

**Figure 4** | CCMs fail to form in most germ-free mice. a, Experimental design in which offspring of susceptible Krit1ECKO females were fostered to conventional or germ-free Swiss Webster mothers. b, Hindbrains from P10 offspring fostered to conventional or germ-free conditions (8 out of 8 shown, top) or germ-free conditions (8 out of 8 shown, bottom). c, Lesion volume quantification of Krit1ECKO hindbrains following C-section/fostering in conventional or germ-free conditions. d, Relative quantification of neonatal gut bacterial load measured by qPCR of bacterial 16S rRNA gene copies. e, Relative quantification of Krit1 mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. f, Relative load measured by qPCR of bacterial 16S rRNA gene copies. g, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. h, Relative load measured by qPCR of bacterial 16S rRNA gene copies. i, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. j, Relative load measured by qPCR of bacterial 16S rRNA gene copies. k, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. l, Relative load measured by qPCR of bacterial 16S rRNA gene copies. m, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. n, Relative load measured by qPCR of bacterial 16S rRNA gene copies. o, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. p, Relative load measured by qPCR of bacterial 16S rRNA gene copies. q, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. r, Relative load measured by qPCR of bacterial 16S rRNA gene copies. s, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. t, Relative load measured by qPCR of bacterial 16S rRNA gene copies. u, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. v, Relative load measured by qPCR of bacterial 16S rRNA gene copies. w, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. x, Relative load measured by qPCR of bacterial 16S rRNA gene copies. y, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. z, Relative load measured by qPCR of bacterial 16S rRNA gene copies. 

**Figure 5** | CCM susceptibility is associated with increased levels of Gram-negative Bacteroidetes s24-7. a–c, Visual and microCT images of hindbrains from susceptible (top) and resistant (bottom) Krit1ECKO and Ccm2ECKO animals and susceptible Krit1ECKO animals fostered to conventional Swiss Webster (SW) mothers. Scale bars, 1 mm. d, e, Quantification of lesion and brain volumes. Error bars shown as s.e.m. and significance determined by pairwise, two-tailed Student’s $t$-test. $****P < 0.0001$; $*$, $P > 0.05$. Note, conventional Swiss Webster data from Figs 4 and 5 are the same experiment.

**Figure 6** | Treatment of Krit1ECKO mice with TAK-242 demonstrated an approximately 80% reduction in CCM lesion volume. a, b, LPS-induced leakage in the hindbrains from susceptible (top) and resistant (bottom) Krit1ECKO mice. c, Hindbrains from susceptible (top) and resistant (bottom) Krit1ECKO mice. d, Relative load measured by qPCR of bacterial 16S rRNA gene copies. e, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. f, Relative load measured by qPCR of bacterial 16S rRNA gene copies. g, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. h, Relative load measured by qPCR of bacterial 16S rRNA gene copies. i, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. j, Relative load measured by qPCR of bacterial 16S rRNA gene copies. k, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. l, Relative load measured by qPCR of bacterial 16S rRNA gene copies. m, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. n, Relative load measured by qPCR of bacterial 16S rRNA gene copies. o, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. p, Relative load measured by qPCR of bacterial 16S rRNA gene copies. q, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. r, Relative load measured by qPCR of bacterial 16S rRNA gene copies. s, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. t, Relative load measured by qPCR of bacterial 16S rRNA gene copies. u, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. v, Relative load measured by qPCR of bacterial 16S rRNA gene copies. w, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. x, Relative load measured by qPCR of bacterial 16S rRNA gene copies. y, mRNA in lung endothelial cells (LEC) measured by qPCR. Red boxes indicate values for the single germ-free animal with the lowest lesion volume. z, Relative load measured by qPCR of bacterial 16S rRNA gene copies.
bacterial load returning to pre-antibiotic levels (Fig. 6i), except for a single generation 3 animal that developed an intra-abdominal abscess with pronounced splenomegaly (Fig. 6i, red star, Extended Data Fig. 9a). Maternal treatment with vancomycin alone, a broad-spectrum antibiotic specific for Gram-positive bacteria, had no effect on CCM formation, consistent with a causal role for GNB (Extended Data Fig. 9b–h). Measurement of s24-7 levels in the intestines of generation 1, 2 and 3 neonates collected for analysis of lesion volume revealed a significant, sustained reduction in generation 3 relative to generation 1 (Fig. 6k, l), consistent with the observation that resistant Krit1fl/fl and Ccm2fl/fl mothers have lower s24-7 levels than susceptible mothers (Fig. 5h). Conversely, sterile C-section/fostering of resistant Krit1flECKO and Ccm2ECKO pups to conventional Swiss Webster foster mothers with high levels of s24-7 (Fig. 5h) restored CCM susceptibility (Extended Data Fig. 10). These findings provide further evidence that qualitative changes in the bacterial microbiome can alter disease course.

**Discussion**

Designing rational therapies for CCM disease is complicated by the fact that many of the pathogenic events take place within brain endothelial cells of the central nervous system (CNS), where drug delivery is blocked by the blood–brain barrier34. The finding that LPS accelerates CCM formation (Fig. 2a–e) although it is unable to cross the blood–brain barrier35 suggests that CCM formation is driven by activation of endothelial TLR4 receptors on the luminal, blood side of the blood–brain barrier (Fig. 6a). Whether similar pathways exist for GNB/LPS from the gut lumen into circulation (Fig. 6a). TAK-242 or LPS-RS effectively reduce lesion formation, confirming endothelial TLR4 as a ‘druggable’ target for CCM disease (Fig. 6b–d, Extended Data Fig. 8). Existing TLR4 blocking agents developed for sepsis treatment36 could potentially be repurposed as therapies for severe human CCM disease. However, such application will first need to address the requirement for chronic therapy, the potential risk of lethal sepsis, and whether anti-TLR4 therapy will affect existing as well as nascent lesions.

Manipulation of gut microbiome-host interactions is a more exciting potential strategy to treat a life-long disease such as CCM. The microbiome has been associated with many human diseases37, but specific molecular mechanisms by which it contributes to disease pathogenesis have been difficult to define. Our studies support a central role for the gut microbiome and endothelial responses to GNB in the pathogenesis of CCMs. We find that the bacterial microbiome is the primary source of TLR4 ligand required to stimulate CCM formation in mice, and that small qualitative differences in the gut microbiome may have marked effects on the course of CCM disease in this animal model. Although s24-7 is not found in humans, the association of CCM disease susceptibility with this GNB is particularly interesting because it is associated with disruption of the gut epithelial barrier38. Thus, a key step in CCM pathogenesis is predicted to be translocation of bacteria or bacterial LPS from the gut lumen into circulation (Fig. 6a). Whether similar inflammatory/collagenic microorganisms also accelerate human CCM disease remains an important question. The clinical course of CCM disease is exceptionally variable, even among individuals with familial CCM disease due to a common KRIT1 mutation39,40. Genetic polymorphisms that alter TLR4 and CD14 expression account for some of this heterogeneity (Fig. 3), but most of the clinical variability remains unexplained and may reflect the effect of individual microorganisms. Future studies that simultaneously define the genomes and microbiomes of CCM patients will be required to test this intriguing hypothesis and determine whether the microbiome is a viable therapeutic target for this disease.

**Online Content** Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Contributions A.T.T. designed and performed most of the experiments. J.P.C. and X.Z. performed parallel studies in Sydney. J.K. and J.H.-M. performed experiments. A.T.T. designed and performed most of the experiments.
METHODS

University of Pennsylvania mice. The Cdh5(PAC)-CreERT2 transgenic mice (iEcre) were a gift from R. H. Adams39, Krit1Cre and CmCmCreERT2 animals have been previously described40,41. Tlr4−/−, Cd14−/−, Ali14 (R26-LSL-RFP), and R6-CreERT2 animals42-45 were obtained from the Jackson Laboratories. The Sko1c1(BAC)-CreERT2 transgenic mice have been previously described46. All experimental animals were maintained on a mixed 129/Sv, C57BL/6J, DBA/2 genetic background unless specifically described. C57BL/6J and timed pregnant Swiss Webster mice were purchased from Charles River Laboratories. Germ-free Swiss Webster mice were purchased from Taconic. Breeding pairs between two and ten months of age were used to generate the neonatal CCM mouse model pups. Mice were housed in a specific pathogen-free facility where cages were changed on a weekly basis; ventilated cages, bedding, food, and acidified water (pH 2.5–3.0) were autoclaved before use, ambient temperature maintained at 23–25°C, and 5% Cldix-S was used as a disinfectant. Experimental breeding cages were randomly housed on three different racks in the vivarium, and all cages were kept on automatic 12-h light/dark cycles. The University of Pennsylvania Institutional Animal Care and Use Committee (IACUC) approved all animal protocols, and all procedures were performed in accordance with these protocols.

Centenary Institute mice. A group of the resistant CmCmCreERT2 colony was exported to the Centenary Institute, Sydney, Australia, where the mice were permanently maintained as an inbred colony in a quarantine facility. After several generations, this colony uniformly converted to lesion susceptibility. Cages were changed on a weekly basis; ventilated cages, bedding, food and acidified water (pH 2.5–3.0) were autoclaved before use. Ambient temperature was maintained between 22–26°C, and 80% ethanol and F10SC (1:125 dilution of the concentrate, a quaternary ammonium compound) were used as disinfectants. Experimental breeding cages were randomly distributed throughout the vivarium, and all cages were kept on 12-h light/dark cycles. The Sydney Local Health District Animal Welfare Committee approved all animal ethics and protocols. All experiments were conducted under the guidelines/regulations of Centenary Institute and the University of Sydney.

Gnotobiotic animal husbandry. Germ-free Swiss Webster mice were purchased from Taconic and directly transferred into sterile isolators (Class Biologically Clean Ltd) under the care of the Penn Gnotobiotic Mouse Facility. Food, bedding and water (non-acidified) were autoclaved before transfer into the sterile isolators. Ventilated cages were changed weekly, and all cages in the vivarium were kept under 12-h light/dark cycles. Microbiology testing (aerobic and anaerobic culture, 16S nPCR) was performed every ten days and faecal samples were sent to Charles Rivers Laboratories for pathology testing on a quarterly basis. Further details regarding the sterile C-section fostering can be found below. The University of Pennsylvania Institutional Animal Care and Use Committee (IACUC) approved all animal protocols, and all procedures were performed in accordance with these protocols.

Induction of the neonatal CCM mouse model. For all neonatal CCM mouse model experiments, at one day post-birth (P1), pups were intragastrically injected with 30-gauge needle with 40 ng of 4-hydroxymatoxin (4OHT, Sigma Aldrich, H79042) dissolved in 9% ethanol/corn oil (volume/volume) vehicle (30 μl total volume per injection). This solution was freshly prepared from pre-measured, 4OHT powder, for each injection. Before injection, the pup skin was sanitized using ethanol wipes. The P1 time point was defined by checking experimental breeding pairs every morning for new litters. The following morning (P1), pups were injected with 4OHT. All experimental pups were subjected to this induction regimen. For the Tlr4 rescue experiment (Fig. 2), and all lineage-tracing experiments, an additional dose of 40 ng of 4OHT was intragastrically delivered at P2 (P1 + 2, two total doses). Pups were then harvested as previously described47 at the specified time points.

Histology. Tissue samples were fixed in 4% formaldehyde overnight, dehydrated in 100% ethanol, and embedded in paraffin. 5-μm-thick sections were used for haematoxylin and eosin (H&E) and immunohistochemistry (IHC) staining. The following antibodies were used for immunostaining: rat anti-PECAM (1:20, Histobio Tech DIA-310), rabbit anti-pMLC2 (1:200, Cell Signaling 36745), goat anti-KLF4 (1:100, R&D AF3158), and rabbit anti-RFP (1:50, Rockland 600-401-379). Littermate control and experimental animal sections were placed on the same slide and immunostained under identical conditions. Images were taken at the same time using the same exposure times and colour channels, and were subsequently overlaid using ImageJ.

Gram staining. Intra-abdominal abscesses were dissected and triturated in 500 μl of SOC medium. Drops of the mixture were placed on a microscope slide, briefly exposed to heat, and Gram staining was performed using a kit from Sigma Aldrich (77730) following the manufacturer’s protocol.

Whole-mount retinal endothelium staining. Eyes from euthanized P17 mice were removed and fixed overnight in cold 4% PFA/PBS solution. The following day, retinas were dissected, cut into petals, and stained with isoelectric-B4 conjugated to Alexa488 fluorophore (Thermo Fisher 121411) as previously described48. The retinas were then whole-mounted on microscopy slides in a flat, four-petal shape for fluorescence imaging.

Bacteroides fragilis abyssus model. B. fragilis was purchased directly from the ATCC (strain 25285) and grown in chopped meat glucose (CMG) broth (Anaerobe Systems AS-813) under anaerobic conditions at 37°C. Autoclaved, deagassed caecal contents (ACC) were generated by collecting caecal contents from the colons of euthanized adult mice between 2–8 months of age. Caecal contents were then autoclaved and pulverized in an equal volume of CMG broth. This slurry was mixed with ACC and underwent intra-cardiac perfusion in the anaerobic chamber. 1 ml of CMG broth was inoculated with B. fragilis and grown overnight to an optical density of between 0.8 and 1.0. An equal volume of ACC was mixed with the overnight bacterial culture. 100 μl of this B. fragilis-ACC mixture was injected intraperitoneally into five-day-old pups with a 31-gauge needle. Control littersmates were simultaneously injected intraperitoneally with 100 μl of ACC alone. Pups were harvested at P17. Spleen weight was measured immediately after dissection, and all tissue was subsequently processed as described above.

Intravenous LPS, peptidoglycan, poly(I:C), IL-1β and TNFα injections. LPS from E. coli O127:B8 was purchased from Sigma (L3129) and administered to the low-lesion-penetrance, resistant CmCmCreERT2 neonatal CCM disease model. At P5, a 3 μg dose of LPS dissolved in sterile PBS was administered retro-orbitally in a total 30 μl volume by 31-gauge needle. At P10, a 5 μg dose of LPS was administered retro-orbitally in a total 50 μl volume by 31-gauge needle. Control animals were identically injected with PBS alone. Pups were euthanized and brains dissected at specified time points. Peptidoglycan from Bacillus subtilis (a Gram-positive gut commensal) was purchased from Invivogen (tlr1-pgns) and administered to the resistant CmCmCreERT2 neonatal CCM disease model under identical conditions as the LPS experiments. Poly(I:C) was purchased from Invivogen (tlr3-picw) and administered to the resistant CmCmCreERT2 neonatal CCM disease model under identical conditions as the LPS experiments.

Mouse IL-1β was purchased from Genscrit (Z02988) and administered to the resistant CmCmCreERT2 neonatal CCM disease model. At P5, a 5 ng dose of IL-1β dissolved in sterile PBS was administered retro-orbitally in a total 30 μl volume by 31-gauge needle. At P10, an 8 ng dose of IL-1β was administered retro-orbitally in a total 50 μl volume by 31-gauge needle. Control animals were identically injected with PBS alone. Pups were euthanized and brains dissected at specified time points. Mouse TNFα was purchased from Genscrit (Z02918) and administered to the resistant CmCmCreERT2 neonatal CCM disease model under identical conditions as the LPS experiments.

Contrast-enhanced microCT. For all experiments using microCT quantification of CCM lesion volume, brains were harvested and immediately placed in 4% PFA/PBS fixative. Brains remained in fixative until staining with non-destructive, iodine contrast and subsequent microCT imaging performed as previously described47. All tissue processing, imaging and volume quantification were performed in a blinded manner by investigators at the University of Chicago without any knowledge of experimental details.

We blinded samples at three distinct points in the analysis. First, neonatal CCM disease model pups were injected with 4OHT without knowledge of genotypes. Second, hindbrains from mice were given randomized, de-identified labels to provide for blinded microCT scanning by an independent operator. Third, randomized microCT image stacks were analysed in a blinded manner by individuals not involved in any prior experimental steps.

Immune cell isolation from neonatal brain. Mice were anaesthetized with Avertin and underwent intra-cardic perfusion with 10 ml of cold PBS. The brain was separated from the brainstem, and the cerebellum was separated from the remaining brain and processed in parallel. The tissue was minced with scissors, placed in digestion buffer (RPMI, 20 mM HEPES, 10% FCS, 1 mM CaCl2, 1 mM MgCl2, 0.05 mg ml−1 Liberase (Sigma), 0.02 mg ml−1 Dnase I (Sigma), and 1 mM PMSF), filtered through a 70-μm cell-strainer, and diluted to 100–150 μl volume by 31-gauge needle. At P10, a 5 μg dose of LPS was administered retro-orbitally in a total 50 μl volume by 31-gauge needle. Control animals were identically injected with PBS alone. Pups were euthanized and brains dissected at specified time points. Mouse TNFα was purchased from Genscrit (Z02918) and administered to the resistant CmCmCreERT2 neonatal CCM disease model under identical conditions as the LPS experiments.

Haematopoietic cell isolation from neonatal whole blood, spleen and subsequent FACS analysis. Neonatal P10 mice were anaesthetized with Avertin and underwent intra-cardic puncture/blood draw using a 27-gauge needle/syringe coated with 0.5 mM EDTA, pH 8.0 immediately before use. Cells were pelleted at 300 g for 5 min at 4°C. Serum was removed and red blood cells were lysed using ACK lysis buffer. Splenocytes were disected in parallel, hand-homogenized using a mini-pestle and red blood cells were lysed using ACK lysis buffer. Cells from both sets of tissues were passed through a 70-μm cell-strainer, pelleted and resuspended in FACS buffer (PBS, 2% FBS, 0.1% NaN3) for immunostaining and subsequent FACS analysis.

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Immune cell staining and flow cytometry analysis. Cells were isolated from the indicated tissues. Single-cell suspensions were stained with CD16/32 and with indicated fluorochrome-conjugated antibodies. Live/Dead Fixable Violet Cell Stain (Invitrogen) was used to exclude non-viable cells. Multi-laser, flow cytometry analysis procedures were performed at the University of Pennsylvania Flow Cytometry and Cell Sorting Facility using BD LSRII cell analysers running FACSDiva software (BD Biosciences). Two-laser, flow cytometry analyses were performed at the University of Pennsylvania iPS Cell Core using BD Accuri C6 instrument (BD Biosciences). Genotypes were input by Impute v2 (ref. 51) using the GIANT 100G v5 integrated call set for all ancestries as a reference.2 Gene expression levels were measured by Illumina HT12v3 arrays. Gene expression pre-processing involved quantile normalization, log transformation, probe centring and scaling, population stratification correction (first four genetic multi-dimensional scaling components were removed from gene expression data) and correction for unknown confounders (first 20 gene expression principal components not associated with genetic variants were removed from gene expression data). Identification of potential sample mix-ups was conducted by MixupMapper21 and finally 1,227 samples remained. All pre-processing steps were performed with the QTL mapping pipeline v1.2.4D (https://github.com/molgenis/systemsgenetics/tree/master/eqtl-mapping-pipeline - downloading-the-software).

These results are corroborated by an independently conducted GTEx Consortium study (http://www.gtexportal.org/home/snp/rs10759930 and http://www.gtexportal.org/home/snp/rs778587). TAK-242 and LPS-ARDS administration. TAK-242 was purchased from EMD Millipore (614316) and administered to the neonatal CCD disease model. Five, seven and nine days after birth, a 60-μg dose of TAK-242 was dissolved in DMSO/sterile intralipid (Sigma, 1141) vehicle and administered retro-orbitally in a total volume of 30μl. Control animals were identically injected with sterile DMSO/sterile intralipid vehicle alone. Pups were euthanized and brains dissected 10 days after birth. LPS-ARDS ultrapure was purchased from Invivogen (Irl-pulsps) and administered to the neonatal CCD disease model. Starting at P5, a 20μg dose dissolved in sterile PBS was administered retro-orbitally in a total volume of 30μl every 24h. Control animals were identically injected with sterile PBS alone. Pups were euthanized and brains dissected 10 days after birth.

Transgenetic antibiotic administration. Experimental breeding pairs of mice, yielding susceptible neonatal CCD pups, were identified by induction of a neonatal CCD litter and evaluation of lesion burden. These breeding pairs then underwent timed matings and at E14.5, both male and female adult mice received antibiotic-laced drinking water mixed with 40 g·l⁻¹ of sucralose and red food colouring. Antibiotic water was replaced daily. The following antibiotics were mixed with 0.22-μm filtered water: penicillin (500 mg·l⁻¹), neomycin (500 mg·l⁻¹), streptomycin (500 mg·l⁻¹), metronidazole (1 g·l⁻¹) and vancomycin (1 g·l⁻¹). Antibiotics were purchased from the Hospital of the University of Pennsylvania pharmacy. The neonatal CCD model was induced as described above. At P10, pups were euthanized and antibiotic water switched to normal drinking water. Experimental breeding pairs were then mated to obtain third generation, post-antibiotic pups.

Vancomycin mono-antibiotic administration. Co-housed, susceptible Krt14^+/− males underwent evening–morning timed matings with a single susceptible Krt14^+/− female. Upon detection of a plug in the morning, the females were subsequently separated into singly-housed cages. At E14.5, female mice were received either vancomycin (1 g·l⁻¹)–laced or untreated (vehicle) sterile-filtered drinking water, changed daily. The drinking water was further mixed with 40 g·l⁻¹ sucralose and red food colouring. Pups were harvested at P11.

Bacterial DNA extraction from neonatal mouse guts and bacterial ribosomal DNA qPCR. The entire neonatal gut was dissected, snap-frozen on dry ice, and stored at −80°C. Bacterial DNA was extracted using the QIAamp DNA Stool Mini Kit (Qiagen 51514 or 51516) and was used to extract bacterial DNA from the neonatal gut. Before commencing the standard QIAGEN protocol, the frozen gut was mixed in the included stool lysis buffer and homogenized with a 5 mm stainless steel bead in a TissueLyser LT (Qiagen 69980) at 50 Hz for 10 min at 4°C. Concentration of the extracted DNA was equalized and 16 ng of DNA was used per qPCR reaction with universal bacterial 16S rRNA gene primers22, two different sets of previously characterized Bacteroidetes q4−7 primers23,24, and Firmicutes primers25. Universal 16S rRNA forward: 5′-ACTGAGAATACGGCACA-3′; universal 16S rRNA reverse: 5′-ATACCGCGCGCCGCTGAC-3′; Bacteroidetes q4−7 RNA set 1 reverse: 5′-CGGCTCAATCTTCTGCGCA-3′; Bacteroidetes q4−7 RNA set 2 forward: 5′-CCACGACCCGCGGTTAATA-3′; Bacteroidetes q4−7 RNA set 2 reverse: 5′-CAGCATCTTGATCTTCTTCT-3′; Firmicutes primers RNA forward: 5′-TCAACACTGAATGACG-3′; Firmicutes primers RNA reverse: 5′-ACCACACCCGAACAC-3′.
Sterile C-section and fostering to conventional Swiss Webster recipient females. Evening–morning timed matings to generate donor susceptible or resistant females yielding Krit1ECKO or Ccm2ECKO pups were performed and time-pregnant Swiss Webster females (Charles River 024) served as foster mothers. To prevent delivery of the pups, at E16.5, donor females were injected subcutaneously with 100 μl of a 15 mg ml⁻¹ solution of medroxyprogesterone (Sigma Aldrich, M1629) dissolved in DMSO. The morning of E19.5, the donor mother was euthanized by cervical dislocation and submerged in a warm sterile solution of 1% Virkon/S/PBS (weight/volume) for one minute. The uterus was then incubated at a constant low temperature, submerged in a warm sterile solution of 1% Virkon/S/PBS for one minute and quickly rinsed with warm sterile PBS. Pups were then removed from the uterus and fostered to the Swiss Webster recipient female. The following morning, induction of the neonatal CCM model was performed as described above.

Sterile C-section and fostering to germ-free Swiss Webster recipient females. Timed matings were performed using germ-free Swiss Webster mice housed in sterile isolators under care of the University of Pennsylvania Gnotobiotic Mouse Facility. Simultaneous evening–morning timed matings were also performed using co-housed, susceptible Krit1fl/fl females and Krit1fl/fl males previously characterized to yield CCM-susceptible pups. Medroxyprogesterone was administered to donor females and the sterile C-section was performed at E19.5 as described in the previous section. The intact uterus was passed through a 1-tube filled with warm 1% Virkon/S/PBS that was hermetically sealed to the sterile isolator. Pups were dissected from the uterus inside the sterile isolator and fostered to the recipient germ-free Swiss Webster mother. Approximately one week later, fetal samples were collected for microbiology testing. Germ-free status was further confirmed by 16S qPCR of bacterial DNA isolated from maternal faeces and neonatal guts.

Collection of maternal CCM mouse faeces. Fresh faecal pellets were collected from experimental females yielding susceptible or resistant pups one day after harvesting the pups to determine one phenotype severity. Collection was performed between 16:00 and 18:00, pellets were immediately snap-frozen on dry ice, and stored at −80 °C.

Extraction and library preparation of bacterial DNA for 16S rRNA gene sequencing. DNA was extracted from stool samples using the Power Soil htp kit (Mo Bio Laboratories) following the manufacturer’s protocol. Library preparation was performed by using previously described barcoded primers targeting the V1/ V2 region of the 16S rRNA gene55. PCR reactions were performed in quadruplicate using Accuprime Taq DNA Polymerase High Fidelity (Invitrogen). Each PCR reaction consisted of 0.4 μM primers, 1x Accuprime Buffer II, 1 U Taq, and 25 ng of DNA. PCRs were run using the following parameters: 95 °C for 5 min; 20 cycles of 95 °C for 30 s, 56 °C for 30 s, and 72 °C for 90 s; and 72 °C for 8 min. Quadruplicate PCR reactions were pooled and products were purified using AMPureXP beads (Beckman-Coulter). Equimolar amounts from each sample were pooled to produce the final library. Positive and negative controls were carried through the amplification, purification and pooling procedures. Negative controls were used to assess reagent contamination and consisted of extraction blanks and DNA-free water. Positive controls were used to assess amplification and sequencing quality and consisted of gBlock DNA (Integrated DNA Technologies) containing non-bacterial 16S rRNA gene sequences flanked by bacterial V1 and V2 primer binding sites. Paired-end 2 x 250 bp sequence reads were obtained from the MiSeq (illumina) using the 500 cycle v2 kit (Illumina).

Analysis of 16S sequencing. Sequence data were processed using QIIME version 1.9.1 (ref. 58). Read pairs were joined to form a complete V1/V2 amplicon sequence. Resulting sequences were quality filtered and demultiplexed. Operational taxonomic units (OTUs) were selected by clustering reads at 97% sequence similarity56. Taxonomy was assigned to each OTU with a 90% sequence similarity threshold using the Greengenes reference database57. A phylogenetic tree was inferred from the OTU data using FastTree39. The phylogenetic tree was then used to annotate and unweight unweighted UniFrac distances between various samples in the study46,58. Microbiome compositional differences were visualized using principal coordinate analysis (PCoA). Community-level differences between mice genetic background as well as disease susceptibility groups were assessed using a PERMANOVA test44 of weighted and unweighted UniFrac distances. To assess significance in the PERMANOVA test, each cage was randomly re-assigned to groups 9,999 times. Differential abundance was assessed for taxa present in at least 80% of the samples, using generalized linear mixed-effects models. For tests of taxon abundance, the cage was modeled as a random effect, as previous research revealed that faecal microbiota of mice are correlated within cages48. The P-values were corrected for multiple testing using Benjamini–Hochberg method. Statistics. Sample sizes were estimated on the basis of our previous experience with the neonatal CCM model and lesion volume quantification by microCT9. Using 40 historically collected, susceptible Krit1ECKO and Ccm2ECKO P10 brains, we calculated a sample standard deviation of 0.250 mm². Between Krit1ECKO and Ccm2ECKO genotypes, an F-test to compare variances confirmed no significant difference (P = 0.340). Thus, for a two-group comparison of lesion volumes, each sample group requires seven animals for a desired statistical power of 95% (β = 0.05), and a conventional significance threshold of 5% (α = 0.05) assuming an effect size of 50% (0.5) and equal standard deviations between sample groups. These predictive calculations were corroborated by our recent publication in which larger effect sizes (>90%) were found to be statistically significant with four to five samples per group9. All experimental and control animals were littermates and none were excluded from analysis at the time of harvest. Experimental animals were lost or excluded at pre-defined points: (i) failure to properly inject HOFT and observation of significant leakage; (ii) death before P10 because of infection or chaos. Given the early time points, no attempt was made to distinguish or segregate results based on neonatal genders. P values were calculated as indicated in figure legends using an unpaired, two-tailed Student’s t-test; one-way ANOVA with multiple comparison corrections (Holm–Sidak or Bonferroni); PERMANOVA; or linear mixed effects modelling. As indicated in the figure legends, the standard error of the mean (s.e.m.), 95% confidence interval, or boxplot is shown.

Data availability. All relevant data are available from the authors upon reasonable request.
Extended Data Figure 1 | CCM formation in resistant Ccm2\textsuperscript{ECKO} animals is stimulated by abscess formation and LPS. a, Resistance to CCM formation is maintained in a C57BL/6j strain background. Ccm2\textsuperscript{ECKO} (iECre;Ccm2\textsuperscript{fl/fl}) animals were backcrossed seven generations onto a C57BL/6j background and gene deletion was induced at P1 with visual hindbrain assessment at P10. \( n = 7 \). Scale bars, 1 mm. b, Retinal CCM formation is stimulated by GNB infection. Retinas of P17 resistant Ccm2\textsuperscript{ECKO} littermates are shown. The sample shown below is from an animal that developed the spontaneous Gram-negative abscess shown in Fig. 1c. Scale bars, 500 \( \mu m \). c, d, Administration of LPS does not drive CCM formation in Cre-negative neonatal mice. LPS was administered intravenously to Ccm2\textsuperscript{fl/fl} and Ccm2\textsuperscript{ECKO} littermates as shown in Fig. 1g. and hindbrains assessed at P17 visually (c) and histologically (H&E staining; d). \( n \geq 3 \) per group. Scale bars, 1 mm (c) and 100\( \mu m \) (d). e, LPS induces myosin light chain activation in CCM-deficient brain endothelial cells. Phospho-myosin light chain (pMLC) and PECAM staining of hindbrains from P5 LPS- or vehicle-injected resistant Ccm2\textsuperscript{ECKO} littermates. Dotted lines trace the Purkinje cell layer. \( n \geq 4 \) per group. Scale bars, 50\( \mu m \). f, Tlr4 expression does not differ between CCM susceptible and resistant animals. Tlr4 expression was measured using qPCR in cerebellar endothelial cells isolated from the indicated animals at P10. Error bars shown as s.e.m. and significance determined by unpaired, two-tailed Student’s \( t \)-test. n.s., \( P > 0.05 \).
Extended Data Figure 2 | See next page for caption.
Extended Data Figure 2 | Analysis of immune cells in P6 and P11 Krit1<sup>fl/fl</sup> and Krit1<sup>ECKO</sup> brains. a, Gating strategy for B cells, natural killer (NK) cells, γδ T cells, CD4 T cells, CD8 T cells, eosinophils, neutrophils and monocytes/macrophages from cerebrum and cerebellum is shown. Cellular surface markers used were as follows: neutrophils (CD45<sup>+</sup>, CD11b<sup>+</sup>, Ly6-G<sup>+</sup>), eosinophils (CD45<sup>+</sup>, CD11b<sup>+</sup>, CD11c<sup>−</sup>, Ly6G<sup>−</sup>, Siglec-F<sup>+</sup>, SSC<sup>hi</sup>), monocyte/macrophage (CD45<sup>+</sup>, CD11b<sup>+</sup>, CD11c<sup>−</sup>, Ly6G<sup>−</sup>, Siglec-F<sup>−</sup>, SSC<sup>lo</sup>), NK cells (CD45<sup>+</sup>, CD11b<sup>−</sup>, CD19<sup>−</sup>, NK1.1<sup>+</sup>), B cells (CD45<sup>+</sup>, CD11b<sup>−</sup>, CD19<sup>−</sup>, CD19<sup>+</sup>, NK1.1<sup>−</sup>, CD19<sup>+</sup>), γδ T cell (CD45<sup>+</sup>, CD11b<sup>−</sup>, CD19<sup>−</sup>, CD3<sup>+</sup>, TCR<sup>γδ</sup>), CD4 T cell (CD45<sup>+</sup>, CD11b<sup>−</sup>, CD19<sup>−</sup>, CD3<sup>+</sup>, CD19<sup>−</sup>, CD3<sup>+</sup>, TCR<sup>γδ</sup>, CD4<sup>+</sup>, CD4<sup>+</sup>), CD8 T cell (CD45<sup>+</sup>, CD11b<sup>−</sup>, CD19<sup>−</sup>, CD3<sup>+</sup>, TCR<sup>γδ</sup>, CD4<sup>+</sup>). b, The number of B cells, NK cells, γδ T cells, CD4 T cells, CD8 T cells, eosinophils, neutrophils and monocytes/macrophages isolated from P6 cerebrum (top) and cerebellum (bottom) is shown for susceptible Krit1<sup>fl/fl</sup> and Krit1<sup>ECKO</sup> littermates. n ≥ 6 per group. No significant differences were detected. c, The number of B cells, NK cells, γδ T cells, CD4 T cells, CD8 T cells, eosinophils, neutrophils and monocytes/macrophages isolated from P11 cerebrum (top) and cerebellum (bottom) is shown for susceptible Krit1<sup>fl/fl</sup> and Krit1<sup>ECKO</sup> littermates. n ≥ 6 per group. d, e, Frequency of RORγt<sup>+</sup> CD4 T cells isolated from P6 and P11 cerebellum. n ≥ 6 per group. Error bars of all graphs shown as s.e.m. and significance determined by unpaired, two-tailed Student's t-test. *P < 0.05. Note that there is significant immune cell presence in the cerebellum of susceptible Krit1<sup>ECKO</sup> animals at P11 but not at P6.
Extended Data Figure 3 | Changes in the volume of CCM lesions are not accompanied by changes in total brain volume. The indicated total brain volumes were measured using microCT imaging. a, b, Brain volumes corresponding to the genetic rescue experiments shown in Fig. 2c–f, respectively. c, Brain volumes corresponding to the C-section/germ-free fostering experiment shown in Fig. 4b, c. d, Brain volumes corresponding to the intergenerational antibiotic experiment shown in Fig. 6f–h.

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Extended Data Figure 4 | Lineage tracing of the Cdh5(PAC)-CreERT2 transgene in neonatal mice. a–c, R26-LSL-RFP, R26-CreERT2, R26-LSL-RFP, and Cdh5(PAC)-CreERT2, R26-LSL-RFP neonates were induced with doses of tamoxifen on P1+2 (two total doses) and CD45⁺ haematopoietic cell numbers in the spleen and peripheral blood were assessed at P10. n ≥ 5 per group. Error bars shown as s.e.m. and significance determined by one-way ANOVA with Holm–Sidak correction for multiple comparisons. ***P < 0.001; n.s., P > 0.05. Note, the number of labelled haematopoietic cells in Cdh5(PAC)-CreERT2, R26-LSL-RFP animals is indistinguishable from R26-LSL-RFP negative control animals, whereas >90% of CD45⁺ cells were RFP⁺ in R26-CreERT2, R26-LSL-RFP positive control animals. d, Anti-RFP and anti-PECAM immunostaining of P10 hindbrains from Krit1fl/fl-R26-LSL-RFP-negative control and Krit1ECKO-R26-LSL-RFP was performed to identify Cre⁺ descendants at the site of CCM formation. Note that all RFP⁺ cells in Krit1ECKO-R26-LSL-RFP animals are PECAM⁺, consistent with endothelial-specific Cre activity. Asterisk indicates CCM lesion. Results are representative of ≥3 per group. Scale bars, 100 μm.
Extended Data Figure 5 | The Slco1c1(BAC)-CreERT2 transgene is selectively expressed in brain endothelial cells and confers CCM formation when used to drive deletion of Krit1 in neonatal mice. a, R26-LSL-RFP, Cdh5(PAC)-CreERT2-R26-LSL-RFP and Slco1c1(BAC)-CreERT2-R26-LSL-RFP neonates were induced with tamoxifen injection on P1 + 2 (two total doses). Immunostaining for RFP and PECAM was performed at P10 in the indicated tissues. Results are representative of at least three animals per group and three independent experiments. Scale bars, 100 μm. Note the presence of RFP + PECAM + cells in the brain, small intestine, caecum, colon and liver of Cdh5(PAC)-CreERT2-R26-LSL-RFP animals, but only in the brain of Slco1c1(BAC)-CreERT2-R26-LSL-RFP animals. b, Visual (top) and corresponding microCT (bottom) images of brains from susceptible Slco1c1(BAC)-CreERT2-Krit1fl/+, and Slco1c1(BAC)-CreERT2-Krit1fl/fl P10 animals. Arrow indicates CCM lesions in the cerebrum. Scale bars, 1 mm. c, H&E staining of cerebellum (hindbrain) from the indicated animals (left). H&E staining of cerebrum (forebrain) from the indicated animals (middle). KLF4 and PECAM immunostaining from the indicated animals (right). Scale bars, 50 μm. Asterisks denote CCM lesions. n ≥ 5 per group.
Extended Data Figure 6 | CCM formation can be stimulated by IL-1β or poly(I:C) treatment. a, Schematic of the experimental design in which littermates receive a retro-orbital injection of the indicated cytokine or TLR ligand at P5 and P10 before tissue harvest and analysis at P17. b–m, Visual images and volumetric quantification of CCM lesions in the hindbrains of P17 Ccm2<sup>−/−</sup> littermates injected with the indicated cytokines, TLR ligands, or vehicle control are shown. Error bars shown as s.e.m. and significance determined by unpaired, two-tailed Student’s t-test. *P < 0.05; n.s., P > 0.05. Scale bars, 1 mm.
Extended Data Figure 7 | 16S rRNA sequencing results from susceptible and resistant Krit1<sup>fl/fl</sup> and Ccm2<sup>fl/fl</sup> dams. a, Heat map showing relative abundance of bacterial taxa (right) identified in susceptible (blue) and resistant (salmon) Krit1 (ccm1, purple) and Ccm2 (ccm2, green) animals (top). b, Boxplots of bacterial taxa that demonstrated significant differential abundance in susceptible versus resistant animals and the relative abundance of those taxa. c, Boxplot of the Firmicutes (Ruminococcus) taxon that displayed significant differential abundance between Krit1 and Ccm2 genotypes. Note that the relative abundance of Bacteroidetes s24-7 is anywhere from 10-fold to 10,000-fold greater than any other taxon. Significance ($P < 0.05$) for b and c determined by linear mixed effects modelling with Benjamini–Hochberg correction for multiple comparisons.
Extended Data Figure 8 | Blockade of CCM formation by the TLR4 antagonist LPS-RS. 

**a**, Schematic of the experimental design in which Krit1<sup>ECKO</sup> littermates receive retro-orbital injections of the TLR4 antagonist LPS-RS. 

**b**, Visual (left) and microCT (right) images of hindbrains from vehicle or LPS-RS injected animals. 

**c, d**, Quantification of CCM lesion and brain volume in Krit1<sup>ECKO</sup> littermates treated with vehicle or LPS-RS. Error bars shown as s.e.m. and significance determined by unpaired, two-tailed Student’s t-test. **P** < 0.01; n.s., indicates P > 0.05. All scale bars, 1 mm.
Extended Data Figure 9 | CCM formation is stimulated by spontaneous abscess formation and not blocked by vancomycin.  

**a.** P10 hindbrains from generation 3/post-ABX *Krit1*ECKO littermates in the longitudinal antibiotic experiment described in Fig. 6e–l. The animal with a large CCM lesion burden on the far right was found to have an abdominal abscess (circle, 'absc') and splenomegaly (arrow, lower right). Scale bar, 1 mm.  

**b.** Schematic of the experimental design in which cohoused, lesion susceptible *Krit1*ECKO mating pairs were used to test the acute effect of vancomycin treatment on CCM formation. Offspring were studied after receiving maternal vehicle or vancomycin administered from E14.5 to P11.  

**c, d.** Visual images of hindbrains from representative offspring following vehicle or vancomycin antibiotic treatment. Scale bars, 1 mm.  

**e, f.** Volumetric quantification of CCM lesions and brain volumes in *Krit1*ECKO littermates treated with vehicle or vancomycin.  

**g, h.** Relative quantification of total neonatal gut bacterial load measured by qPCR of bacterial universal 16S or Firmicutes-specific rRNA gene copies. n ≥ 6 per group. Error bars of all graphs shown as s.e.m. and significance determined by unpaired, two-tailed Student’s *t*-test. n.s., *P* > 0.05. **∗∗∗∗** *P* < 0.0001.
Extended Data Figure 10 | CCM formation is conferred to the offspring of resistant animals by fostering to Swiss Webster mothers. a, Schematic of the experimental design in which timed matings of resistant Krit1ECKO and resistant Ccm2ECKO mating pairs were used to generate E19.5 offspring delivered by natural birth and raised by the birth mother or C-section/fostered to conventional Swiss Webster foster mothers. b, c, Visual images of hindbrains from P10 resistant Krit1ECKO and Ccm2ECKO offspring following natural delivery and nursing by resistant mothers or after C-section/fostering to Swiss Webster mothers. n ≥ 6 per group.