Conditional deletion of Ccm2 causes hemorrhage in the adult brain: a mouse model of human cerebral cavernous malformations

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Cerebral cavernous malformations (CCM) are irregularly shaped and enlarged capillaries in the brain that are prone to hemorrhage, resulting in headaches, seizures, strokes and even death in patients. The disease affects up to 0.5% of the population and the inherited form has been linked to mutations in one of three genetic loci, CCM1, CCM2 and CCM3. To understand the pathophysiology underlying the vascular lesions in CCM, it is critical to develop a reproducible mouse genetic model of this disease. Here, we report that limited conditional ablation of Ccm2 in young adult mice induces observable neurological dysfunction and reproducibly results in brain hemorrhages whose appearance is highly reminiscent of the lesions observed in human CCM patients. We first demonstrate that conventional or endothelial-specific deletion of Ccm2 leads to fatal cardiovascular defects during embryogenesis, including insufficient vascular lumen formation as well as defective arteriogenesis and heart malformation. These findings confirm and extend prior studies. We then demonstrate that the inducible deletion of Ccm2 in adult mice recapitulates the CCM-like brain lesions in humans; the lesions display disrupted vascular lumens, enlarged capillary cavities, loss of proper neuro-vascular associations and an inflammatory reaction. The CCM lesions also exhibit damaged neuronal architecture, the likely cause of neurologic defects, such as ataxia and seizure. These mice represent the first CCM2 animal model for CCM and should provide the means to elucidate disease mechanisms and evaluate therapeutic strategies for human CCM.

INTRODUCTION

Cerebral cavernous malformations (CCM) are vascular lesions that occur most commonly in the central nervous system of patients (reviewed in 1,2). The lesions are prone to hemorrhage; they are histologically characterized by thin-walled, enlarged capillary cavities, lacking smooth muscle support. Symptoms in patients range from severe headaches to seizures, neurologic disorders and ultimately strokes that can lead to death. CCM lesions are detected at a frequency of 1 in 200–250 individuals in the general population and they occur in a sporadic or in a familial form. The familial form is inherited in an autosomal dominant fashion with incomplete penetrance (1). The disease is associated with inherited loss-of-function mutations in one allele of three distinct genes, CCM1 [KRIT1 (KREV1/RAP1A interaction trapped-1)], CCM2 [OSM (Osmosensing scaffold for MEKK3): Malcavernin; MCG4607] or CCM3 [PDCD10 (Programmed cell death 10)] (1,2). Many germline mutations have been identified, and with few exceptions, they appear to severely cripple or fully inactivate the expression of the encoded protein. In recent years, evidence has mounted implicating a somatic loss of the second allele as the ultimate cause of the disease. Although technologically challenging, careful

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analyses of lesions have uncovered second mutations in the allele opposite the inherited one in lesions, which would completely eliminate expression of a functional CCM protein from both alleles of the affected locus, at least in a portion of the cells within a given lesion (3–6).

Critical insight into the in vivo functions of CCM proteins was gained from mouse models genetically engineered to lack expression of CCM proteins (1). Loss of either Ccm1 (7), Ccm2 (8–10) or, most recently, Ccm3 (11) resulted in embryonic lethality, associated with grossly defective angiogenesis. The targeted null mutations in Ccm genes caused growth arrest after E8.5 and the embryos were generally dead by E10–11. Growth arrest was accompanied by defects in angiogenic remodeling in the peripheral vasculature and lumen formation in the branchial arch artery connecting the heart with the dorsal aorta, thus apparently constituting blood flow and resulting in failure to fully establish circulation. These data seem to suggest that CCMs are required for proper angiogenesis; however, the potential requirement for blood flow in angiogenesis makes it difficult to distinguish a primary requirement of CCM proteins in angiogenesis from a secondary defect due to abnormal heart development and function. Furthermore, the early embryonic lethality in mice with homozygous knockout of Ccm genes precludes studies of the role of CCM in the cerebral vasculature.

Whether human CCM lesions are caused by loss of CCM genes in vascular cells or neuronal cells has been a matter of debate (12,13). More recently, conditional loss of Ccm genes in vascular endothelial cells (ECs), smooth muscle cells (SMCs) and/or neuronal cells in mice has been accomplished (8–11). Deletion of floxed Ccm2 or Ccm3 in ECs using a Tie2-Cre driver caused cardiovascular defects, resembling the phenotypes seen in conventional knockout of Ccm2 or Ccm3, whereas loss of Ccm2 or Ccm3 in neuronal cells using a Nestin-Cre driver or loss of Ccm3 in SMCs using a SM22α-Cre driver failed to replicate such phenotypes. These findings do indicate that expression of CCM proteins in ECs is required for proper cardiovascular development, although a very recent study suggests that loss of Ccm3 in at least some neuronal cells may also have a role in forming proper vascular structures (14). Several additional lines of evidence have suggested roles for CCM proteins in cellular processes of particular relevance to endothelial cell functions. A prior ultrastructural analysis of excised CCM lesions from processes of particular relevance to endothelial cell functions.

**RESULTS**

**Generation of Ccm2 mutant mice**

CCM2 was initially cloned via yeast-two-hybrid screens as an adaptin protein (termed OSM) for the ‘Mitogen-Activated Protein Kinase Kinase Kinase 3’ (MEKK3), a kinase capable of activating the transcription factor NF-κB (18–20) (unpublished data). Subsequently this gene was identified as one of the three genetic loci carrying inactivating mutations in one of its alleles in patients with the inherited form of the CCM disease (12,21). To study the function of Ccm2 in vivo, we first generated conventional Ccm2 knockout mice (Supplementary Material, Fig. S1A). To explore the role of CCM2 specifically in the vasculature during development, but also to explore CCM2’s role in adults, we further generated conditional Ccm2 knockout mice (Supplementary Material, Fig. S1B). On a C57BL/6J background, Ccm2<sup>+/−</sup> heterozygous mice, Ccm2<sup>flox/+</sup> heterozygous mice and Ccm2<sup>flox/flox</sup> homozygous mice are all viable and fertile.

Ccm2<sup>−/−</sup> homozygous mice were embryonic lethal and displayed cardiovascular defects, as previously reported (8–10). By E8.5–E9.5, homozygous embryos were clearly distinguishable from their wild-type or heterozygous littermates due to pericardial edema, a truncated forehead region, growth retardation and arrested heart development (Supplementary Material, Fig. S2). All mutant embryos observed were dead by E10.

**Ccm2 is required for proper angiogenesis**

The yolk sac vasculature is a prominent site of angiogenesis in the early embryo at E8.5–E9.5, where the primitive capillary vascular plexus rapidly remolds to form a well-defined, hierarchical branching network (Fig. 1A, arrowheads). E9.5 Ccm2 homozygous mutant yolk sac displayed a complete lack of remodeled vessels, although the mutants appeared to retain the capillary plexus resulting from vasculogenesis (Fig. 1A). Further analysis of peripheral vasculature and heart development was carried out in the Ccm2 mutants and control littermates (heterozygous embryos) by whole-mount staining for PECAM-1, a pan-endothelial cell marker. In the Ccm2 mutant embryos, the head and intersomitic vasculature failed to assemble properly (Fig. 1B versus C and D versus E).
By E9.5, control littermates had developed the fine branches of the head vasculature (Fig. 1B). In contrast, while the initial head capillary plexus had also formed in the homozygous mutants, it subsequently remained in a primitive state (Fig. 1C). The intersomitic vessels arise from branches of the dorsal aorta and posterior cardinal veins, and grow between adjacent somites to form an intricate network (Fig. 1D). In the homozygous mutants, the vessels appeared to fuse with little branching (Fig. 1E). These phenotypes indicate an angiogenic arrest during cardiovascular development, consistent with previous reports (8–10).

To explore whether endothelial CCM2 is required for the vascular development, we sought to conditionally knock out Ccm2 in ECs. Deletion of Ccm2 in ECs was accomplished with a Tie2-Cre driver, which is active in the endothelium (22). We observed growth arrest in Ccm2flox/flox;Tie2-Cre embryos compared with control littermates, although the arrest was less severe than that of conventional CCM2

Figure 1. Cardiovascular defects in Ccm2 mutant embryos. (A) Yolk sac angiogenesis is defective in Ccm2 mutant embryos. Compared with the developing hierarchical vascular networks with remodeled vessels in the control Ccm2+/+ yolk sac (arrowheads), the Ccm2 mutant yolk sac reveals defective angiogenic remodeling. (B and C) Defective head angiogenesis in Ccm2 mutant embryos. Whole-mount staining with a pan-endothelial cell marker, anti-PECAM-1 antibody shows arrested vascular remodeling in the head of Ccm2 mutant embryos (open arrowheads) compared with littermate Ccm2+/+ controls (arrowheads). (D–G) Defective trunk angiogenesis and heart morphogenesis in Ccm2 mutant embryos. The remodeling of the intersomitic vessels seen in Ccm2+/+ controls (D; arrowheads) is defective in Ccm2 mutant embryos (F; open arrowheads). Growth of the ventricular endocardium appears arrested in Ccm2 mutant embryos (G; open arrowhead) compared with controls (F; arrowhead) (red-dotted line enclosed boxes in D and E shown at higher magnification in F and G, respectively). Scale bars are 100 μm.

Figure 2. Defective arteriogenesis and vascular integrity in conventional and endothelial-specific Ccm2 mutant embryos. (A–F) Triple immunofluorescence confocal microscopy using antibodies to the arterial markers NP1 (A–F, green), together with PECAM-1 (A–C, blue) and smooth muscle cell marker αSMA (A–C, red), as indicated. Both conventional mutant Ccm2−/− and endothelial-specific mutant Ccm2flox/flox; Tie2-Cre embryos exhibit defective arterial differentiation when compared with controls (Ccm2+/fflox; Tie2-Cre) (A versus B and C, D versus E and F, arrowhead versus open arrowhead, respectively). In control littermates, αSMA+ SMCs are associated with the dorsal aorta (A) (red, arrow) but not with vein (V) at E9.5. In contrast, both Ccm2−/− and Ccm2flox/flox; Tie2-Cre mutant embryos lack smooth muscle coverage in dorsal aorta (A versus B and C, red). Sp: spinal cord. Scale bars are 100 μm. (G–O) Triple immunofluorescence confocal microscopy of embryonic dorsal aorta was performed with antibodies to vascular luminal membrane proteins Podocarxyn (PODXL, red) and Moesin (green), together with PECAM-1. In control littermates, the apical surface marker Podocarxyn presents on the vascular lumen and ERM adaptor protein Moesin is localized at the luminal or apical membranes to help form a stable vascular lumen (G, J, M, arrowheads). In contrast, the CCM2 mutant mice do not display a smooth and stable vascular lumen with proper enrichment of Podocarxyn and Moesin (H, I, K, L, N, O, open arrowheads). Between the two CCM2 mutants, the conventional Ccm2−/− mutants display a greater disruption of the luminal surface than the Ccm2flox/flox; Tie2-Cre mutants (H versus I, K versus L and N versus O). Scale bars are 100 μm.
knockouts (data not shown). The head and trunk of such embryos were subjected to whole-mount analysis with anti-PECAM-1 antibody. Ccm2flox/flox;Tie2-Cre embryos failed to properly remodel the vasculature to form a hierarchical branching network in yolk sacs, and head and intersomitic regions (Supplementary Material, Fig. S3A and C versus B and D), identical to what had been observed in conventional Ccm2 homozygous embryos. Taken together, these data suggest that endothelial Ccm2 is essential for embryonic angiogenesis, in agreement with prior reports (8,9).

We further examined the development of the dorsal aorta in the conventional and conditional Ccm2 mutant embryos. By E9.5, dorsal aorta formed with the expression of arterial markers (Fig. 2A and D). The expression of the arterial EC marker NP1 was greatly reduced in the dorsal aorta of both Ccm2 homozygous and Ccm2flox/flox;Tie2-Cre embryos (Fig. 2B, C, E, F). These data suggest that arteriogenesis is deficient in the Ccm2 mutant embryos.

Defective vascular integrity in Ccm2 mutant embryos

The fact that human CCM lesions displayed a disrupted endothelial lumen devoid of smooth muscle coverage that resulted in intracranial hemorrhage prompted us to examine whether the Ccm2 mutant vessels recruit vascular SMCs and form proper endothelial junctions and lumen. By E9.5, dorsal aortae were associated with aSMA+ vascular SMCs in the littermate control embryos (Fig. 2A). In contrast, aSMA+vascular SMCs were not detectable in the aortae of either Ccm2 homozygous or Ccm2flox/flox;Tie2-Cre embryos (Fig. 2B and C). Because endothelial cell-specific deletion of Ccm2 resulted in defective smooth muscle coverage, this suggests that CCM2-deficient ECs were defective in the recruitment of these cells. We next examined the distribution of the luminal surface marker Podocynx (PODXL) and the Ezrin–Radixin–Moesin adaptor protein, both of which localize to endothelial cell–cell contacts and are enriched at the plasma membrane facing the developing vascular lumen (23). We found that the conventional and EC conditional Ccm2 mutant embryos failed to form a proper aortic lumen; deletion of Ccm2 resulted in a disrupted lumen compared with what was observed in control littermates (Fig. 2G versus H; I and J versus K; L and M versus N and O). These results indicate that Ccm2 function is required for proper endothelial lumen formation.

Ccm2 function is required for heart development

We first carefully assessed the cardiac phenotype of Ccm2 mutants by using whole-mount staining with anti-PECAM-1 antibody. Obvious defects in cardiac morphogenesis were seen in both Ccm2 homozygous and Ccm2flox/flox;Tie2-Cre embryos (Fig. 1F versus G and Supplementary Material, Fig. S3E versus F). The cardiac looping was incomplete and the PECAM-1+endocardium failed to expand. These findings were confirmed by analysis of the endocardium in stained sections which revealed that the Ccm2 mutant hearts had no or significantly deficient ventricular trabeculation (Supplementary Material, Fig. S4A versus B and C; D versus E and F).

Conditional deletion of Ccm2 leads to hemorrhage in the adult brain

Since conventional or endothelial-specific deletion of Ccm2 resulted in embryonic lethality, these mice do not represent models for the human disease, which occurs in mid-life and which is characterized by lesions largely confined to capillaries in brains. However, the cardiovascular defects detected in early embryos did prompt us to generate Ccm2flox/flox;MX1-Cre mice, because such mice afforded us the opportunity to conditionally delete the Ccm2 gene in adult tissues, and thus test whether deletions in adults might cause brain hemorrhages as seen in the human CCM disease. Previous studies indicated that brain ECs are an efficient target for the MX1-Cre-mediated inducible knockout system (24,25). In agreement with this, we found that the Cre expression is mainly detected in PECAM-1+ ECs but not GFAP+ glial cells in the brain of polyinosinic–polycytidylic acid (pIpC)-treatment of Ccm2flox/flox;MX1-Cre mice (Supplementary Material, Fig. S5, arrowheads). We treated Ccm2flox/flox;MX1-Cre mice and Ccm2flox/+;MX1-Cre control littermates with three injections of pIpC at 2-day intervals to induce Cre expression in 6–8-week-old mice. We did not observe any apparent neurologic or other defects during the first couple of months, during which we inspected mice regularly. However, between 7 and 8 months of age, three out of six Ccm2flox/flox;MX1-Cre mice suddenly and unexpectedly died, while none of the littermate controls suffered this fate: these events prompted us to again more closely observe the remaining pIpC-treated Ccm2flox/flox;MX1-Cre mice. Within a week, we noted that one of these mice had developed a pronounced ataxia with seizure-like behavior. At this juncture, we euthanized all remaining Ccm2flox/flox;MX1-Cre pIpC-treated mice as well as their littermate controls and performed histological analyses of their brains. Gross inspection of whole brains demonstrated that all flox/flox mutant, pIpC-treated mice exhibited prominent hemorrhages, while none of the littermate controls (flox/+; pIpC-treated) exhibited such lesions (Fig. 3A versus B, dorsal view; C versus D, ventral view). It is thus likely that the sudden deaths of the first three pIpC-treated Ccm2flox/flox;MX1-Cre were due also to hemorrhages and associated neurologic defects.

Further analyses of sections revealed that hemorrhages appeared widespread throughout the brain, with larger hemorrhages visible most often in the cerebrum and cerebellum (Fig. 3F and H, respectively; littermate controls Fig. 3E and G). This mosaic phenotype might reflect either the efficiency of Cre-mediated deletion of Ccm2 or the contribution of CCM2-deficient circulating endothelial progenitors to the brain microvasculature (see Discussion). H&E and Nissl stainings confirmed hemorrhages and also showed damaged neuronal architecture in the cerebrum (H&E, Fig. 4A and C versus B and D; Nissl, Fig. 4E and G versus F and H) and cerebellum (H&E, Supplementary Material, Fig. S6A and C versus B and D; Nissl, Fig. S6E and G versus F and H) of pIpC-treated Ccm2flox/flox;MX1-Cre mouse brains. Consistent with the fact that magnetic resonance imaging detects iron deposits in CCM lesions in patients, we also observed the accumulation of hemosiderin, a breakdown product of erythrocytes, as evidenced by brown granules in the affected mice (Fig. 4D,
arrowheads). Damaged neurons were only seen in the CCM lesion (Fig. 4H, open arrowheads). In contrast, we did not find any neurological defects or hemorrhages in the control littermates of the same age. These findings demonstrate that conditional deletion of Ccm2 in adult mice results in hemorrhages in the adult brain, which likely cause neuronal damage and, in consequence, visible neurologic defects, such as ataxia, and ultimately death.

We further addressed whether the brain lesions underlying hemorrhage have defective vascular integrity; specifically, we investigated lumen formation. We found that normal capillaries present in both pIpC-treated Ccm2fl/fl;MX1-Cre mice and their control littermates had a grossly normal Podocarxyn (PODXL)\(^+\) lumen structure (Fig. 5D and E). In contrast, the lesional, CCM-like vessels showed massive accumulation of Ter119\(^+\) erythrocytes, were grossly dilated and exhibited an impaired and disrupted endothelial lumen structure (Fig. 5B, C, F and G; littermate control 5A); the lesional vessels were apparently prone to rupture, resulting in hemorrhage, as evidenced by erythrocytes outside the vessels (Fig. 5B and G).

In addition, CD11b\(^+\) microglia were recruited to CCM lesions (Fig. 6C versus D), where dilated vessels and damaged vessels without extracellular matrix were present (Fig. 6B; littermate control 6A). This suggests an immune response to the CCM pathology (Fig. 6D). Furthermore, GFAP\(^+\) astrocytes failed to extend their projections to surround capillaries (Fig. 6E versus F). In addition, only very few or no Aquaporin4\(^+\) astrocyte end-feet could be detected in CCM lesions (Supplementary Material, Fig. S7). Instead, GFAP\(^+\) astrocytes formed abnormal cell clusters in the vicinity of the lesional vessels (Fig. 6F, arrowheads). Such findings indicate an impaired blood–brain barrier in the CCM lesions. The observed pathological features of the brain lesions in the pIpC-treated Ccm2fl/fl;MX1-Cre mice appear to be highly similar to those described for lesions in human CCM patients (reviewed in 2,26). We also investigated Ccm2fl/fl;MX1-Cre just 3 weeks after pIpC-treatment and were able to detect lesions already at this early time point, although these
Lesions appeared to be mostly smaller (Supplementary Material, Fig. S8). Taken together, our results demonstrate that conditional deletion of *Ccm2* in adult mice causes CCM-like lesions with hemorrhage in the brain, which can result in significant neurologic defects and ultimately death with advancing age.

**DISCUSSION**

Loss-of-function mutations of *Ccm* genes in humans can lead to lesions in brain capillaries that result in hemorrhages that in turn cause apparent neurologic defects and even death in patients. However, the roles of CCM proteins in brain vasculature and their significance in the disease context have remained poorly understood. In the present study, we first provide genetic evidence that endothelial and/or endocardial *Ccm2* expression in mice is required for proper angiogenesis and heart development during embryogenesis, in agreement with previous studies (8–10). We have extended these observations to show that expression of *Ccm2* in ECs is required for formation of smooth and stable vessel lumens, implying a role for CCM2 in endothelial cell–cell interactions. Importantly, we now demonstrate that conditional *Ccm2* gene inactivation in adult mice induces brain hemorrhages highly reminiscent of those observed in human CCM patients. This provides the first CCM2 mouse model for CCMs. The CCM-like brain lesions induced in mice display disrupted vascular lumens and enlarged capillary cavities, lacking neuro-vascular associations; in addition, these lesions are surrounded by accumulations of activated microglia. This CCM animal model should prove to be very valuable to explore the signaling pathways and functions...
of CCM2 and to understand how the absence of CCM2 can lead to severe CCMs in brain capillaries. Furthermore, this animal model may prove useful in testing therapeutic intervention strategies for the human disease.

Our data demonstrate that CCM2 is required in endothelium for the angiogenic remodeling of the vasculature in the yolk sac, the head and intersomitic vessels, but is not required for the initial capillary formation. We also find that expression of arterial markers and recruitment of vascular SMCs are severely impaired in Ccm2 mutants. Previous studies by others and us have indicated that VEGF-A induces arterial differentiation by acting through the Notch pathway (27–29). Furthermore, VEGF-A has also recently been shown to stabilize VEGFR2 via CCM3, thereby facilitating signaling by this receptor (11). Given the ability of CCM2 to associate with both CCM1 and CCM3, CCM2 could conceivably have a role in VEGFR2 signaling as well.

Complete lumen formation in the dorsal aorta is essential for a proper circulatory system. Our extensive confocal imaging analyses, which have made use of several markers for luminal membranes, have revealed that deletion of endothelial Ccm2 resulted in a disrupted lumen in dorsal aorta. This observation suggests that CCM2 may regulate cytoskeletal architecture in ECs. Recent biochemical studies using human microvascular endothelial cells indicate that CCM2 can directly associate with and inhibit RhoA activity, decreasing stress fiber formation; therefore, loss of Ccm2 could lead to disruption of cell–cell junctions and thus lumen formation as a consequence of cytoskeletal changes (9).

During heart development, essential and reciprocal interactions occur between endocardial cells and between endocardial and myocardial cells. Notch has been shown to mediate the endocardium–myocardium interaction that is critical for trabeculation and ventricular chamber morphogenesis (30). Interestingly, endothelial/endocardial deletion of Notch1 or RBPJk results in defective ventricular patterning and trabeculation that is almost identical to the cardiac phenotype seen in the Ccm2 mutants. A recent report also demonstrates that CCM1 may activate Delta–Notch signaling, which leads to AKT phosphorylation in ECs in vitro (31).

The induced loss of Ccm2 in a limited number of blood-accessible cells and ECs in adult mice generated specific, localized lesions in brain capillaries, resulting over time in neurologic defects, hemorrhages and ultimately the death of animals. The lesions observed in the Ccm2 mutants appear to be very similar to those seen in human CCM patients. Mouse brain lesions were characterized by grossly dilated malformed vessels that were prone to rupture, releasing erythrocytes. Furthermore, the normal EC–astrocyte interaction was severely disrupted. These findings indicate defects in EC association of the lesional vessels that are somewhat reminiscent of those seen during angiogenesis in mice lacking CCM2 in ECs during embryonic development. Similar to human lesions, those in brains of adult mice were associated with an immune response as well, as documented by the recruitment of microglia.

How might loss of CCM2 cause lesions in adult brain? Since only a small subset of brain capillaries develops malformations and hemorrhages, we hypothesize that such lesions may develop in newly forming vessels, a process that may be initiated in response to local inflammation or some other event (32,33). If so, one may imagine a scenario, wherein resident Ccm2-deleted ECs and/or EC precursors are recruited into newly sprouting vessels, resulting in a partially defective/disrupted lumen on account of the impaired functions of the Ccm2-deficient ECs. In an alternative and possibly additional scenario, one may imagine that mutant resident ECs and/or their precursors may be prone to initiate new vessel formation and/or preferentially contribute after the process has been initiated, thus increasing the representation of mutant cells in the new vessel wall. In support of this hypothesis, recent reports have suggested that loss of CCM proteins might in fact promote angiogenic sprouting, migration and proliferation of ECs in culture (31,34–36). Conceivably, CCM proteins could have a positive role in transmitting angiogenic signals, while at the same time, setting thresholds to limit spontaneous growth and sprouting.

The brain lesions induced by plpC-treatment of Ccm2flox/flox;MX1-Cre mice over time appear to faithfully recapitulate the pathogenesis of the lesions in human CCM patients. Therefore, this represents the first CCM2 mouse model for the CCM disease. After completion of our work, another mouse model for CCM was recently reported (37). In this model, heterozygous Ccm1+/− mice were crossed onto a background of Msh2 deficiency, a gene critical for mismatch repair. In this model, complete loss of Ccm1 was suggested to occur in some cells as a result of the increased somatic mutational load, resulting in brain lesions resembling those seen in human patients. In contrast, backcrossing of Ccm2+/− mice onto an Msh2 deficient background failed to generate lesions; the reasons for this are unknown. Unlike our model, wherein we induce the loss of Ccm2 in adult mice, the timing of the deletion of Ccm1 in the Msh2-deficient model cannot be controlled and loss of Ccm1 and the generation of lesions could occur at any time starting during embryonic development; furthermore, penetrance in this model was reported at 50% only. Finally, and importantly, Msh2 deficiency likely predisposes towards mutations in other genes, which potentially compromises the usefulness of the model. Nevertheless, the two models are also complementary as they provide a means to compare and contrast the pathogenesis of lesions induced by loss of two different CCM proteins; CCM proteins may have distinct activities in the context of CCMs (36).

The plpC-induced, MX1-Cre-mediated deletion of Ccm2 in adult mice described here represents a temporally controllable and fully penetrant model for generating brain vascular lesions that resemble those observed in human CCM patients. This mouse disease model should greatly facilitate further investigations into the cellular and molecular mechanisms underlying CCM vascular disease and may ultimately allow for testing of possible therapeutic interventions to slow or stop the progression of this disease.

**MATERIALS AND METHODS**

**Mice**

Conventional and conditional Ccm2 knockout mice on the C57BL/6 (B6) background were generated by homologous
recombination in ES cells according to standard procedures (see Supplementary Material, Fig. S1A and B). Knockout founder mice were generated at Ozgene (Ozgene Pty Ltd, Bentley, Western Australia, Australia). Tie2-Cre founder mice were generated at Ozgene (Ozgene Pty Ltd, Bentley, Western Australia, Australia). Tie2-Cre (22) and MX1-Cre (38) mice have been described elsewhere and were crossed with the conditional CCM2 knockout mice. Mice were bred and housed in National Institute of Allergy and Infectious Diseases (NIAID) facilities, and all experiments were done with approval of the NIAID Animal Care and Use Committee and in accordance with all relevant institutional guidelines.

Whole-mount and section immunohistochemistry of mouse embryo
Mouse embryos were dissected between E8.5 and E9.5, fixed in 4% paraformaldehyde/PBS at 4°C for 1–2 h. Staining was performed using anti-PECAM-1 antibody (rat monoclonal antibody, clone MEC 13.3, BD pharmingen, 1:300, overnight at 4°C) to detect endothelial/endocardial cells. HRP-conjugated secondary antibody (Jackson, 1:300, overnight at 4°C) was then used. All brightfield images were captured using QImaging RETIGA 2000R camera (QImaging). For section staining, fixed embryos were sunk in 15% sucrose and 7.5% gelatin in PBS, and then 10 µm sections were collected on a cryostat (Leica). Staining was performed using anti-PECAM-1 antibody (rat monoclonal antibody, clone MEC 13.3, BD pharmingen, 1:300, overnight at 4°C or hamster monoclonal antibody, Chemicon, 1:200, overnight at 4°C) to detect endothelial/endocardial cells, and an anti-NP1 antibody (rabbit polyclonal antibody, A.L. Kolodkin, 1:3000, 3 h at room temperature) as an arterial marker; Cy3-conjugated anti-α SMA antibody to detect SMCs; anti-Moesin antibody (rabbit polyclonal antibody, Abcam, 1:200, overnight at 4°C) and anti-Podocarxyn antibody (rat monoclonal antibody, R&D, 1:200, overnight at 4°C) as vascular lumen markers. TO-PRO-3 (Invitrogen, 1:1000, 10 min at room temperature) was used as a pan-nuclear marker. For immunofluorescent detection, Alexa-488-, Alexa-568-, Cy3-conjugated secondary antibodies (Invitrogen, 1:250, and Jackson, 1:300, 1 h at room temperature) were used. All confocal microscopy was carried out on a Leica SP5 confocal (Leica).

Histology of adult brain
All brightfield images of whole and sectioned brains were captured using QImaging RETIGA 2000R camera (QImaging). For brain-section staining, brains were fixed in 4% paraformaldehyde/PBS at 4°C for overnight, sunk in 15% sucrose in PBS and then 20 µm sections were collected on a cryostat (Leica). Staining was performed using anti-Ter119 antibody (rat monoclonal antibody, BD Serotec, 1:300, overnight at 4°C) to detect erythrocytes, anti-Podocarxyn antibody to detect vascular lumen, anti-collagen type IV antibody; Cy3-conjugated secondary antibodies (Invitrogen, 1:250, and Jackson, 1:300, 1 h at room temperature) were used. All confocal microscopy was carried out on a Leica SP5 confocal (Leica).

plpC administration
Ccm2flox/flox and Ccm2flox/+ mice carrying the MX1-Cre transgene were injected with 250 µg plpC (Sigma-Aldrich, St Louis, MO) three times at 2-day intervals.

SUPPLEMENTARY MATERIAL
Supplementary Material is available at HMG online.

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Conflict of Interest statement. None declared.

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