Arteriovenous Malformations and Other Vascular Malformation Syndromes

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Vascular malformations are a disruption of the normal vascular pattern in which it is expected that a capillary network of microscopic vessels lies interposed between high-pressure arteries that deliver blood and thin-walled veins that collect low-pressure blood for return to the heart. In the case of arteriovenous malformations, arteries or arterioles connect directly to the venous collection system, bypassing any capillary bed. Clinical consequences result from rupture and hemorrhage, from dramatically increased blood flow, or from the loss of capillary functions such as nutrient exchange and filtering function. These malformations can occur sporadically or as a component of inherited vascular malformation syndromes. In these and other hereditary vascular malformation syndromes, genetic studies have identified proteins and pathways involved in vascular morphogenesis and development. A common theme observed is that vascular malformations result from disruption in pathways involved in vascular stability. Here we review the vascular malformations and pathways involved in hereditary hemorrhagic telangiectasia, capillary malformation–arteriovenous malformation, cerebral cavernous malformations, and mucocutaneous venous malformations.

The vertebrate circulatory system depends on the delivery of blood to tissues by arteries and the collection and the return of blood to the heart by veins. In normal circulation, arteries and veins communicate through capillaries, microvascular connections that allow for efficient exchange of nutrients, wastes, oxygen, and carbon dioxide. The control of this system must be dynamic and responsive to changes due to growth, injury, and disease. The complexities involved in tightly regulating this relationship open up opportunities for misregulation and disturbed vascular morphogenesis resulting in vascular malformations.

ARTERIOVENOUS MALFORMATIONS

The normal vascular pattern is disrupted in the case of arteriovenous malformations (AVMs), in which an artery or arteriole connects directly to
a vein or venule, bypassing nutrient exchange and shunting blood directly to the venous system. Arteriovenous malformations consist of clusters of dilated feeding arteries or arterioles that collect to a nidus, where there is a connection to a draining vein or system of veins (Al-Shahi and Warlow 2001). The lesion is termed an “arteriovenous fistula” if there is a connection between a single dilated artery and vein (Al-Shahi and Warlow 2001). AVMs are encountered in both systemic and pulmonary vascular beds and may be isolated lesions or associated with other clinical syndromes. In some organs and tissues, AVMs pose a significant clinical problem. Brain AVMs are a common cause of intracranial hemorrhage, particularly in younger populations (Al-Shahi and Warlow, 2001). AVMs have been described in almost every tissue or organ (Terry et al. 1980; Ishihara et al. 1997; Al-Shahi and Warlow 2001; Matsushige et al. 2004; Buscarini et al. 2005; Cura et al. 2010) and can lead to significant complications. The etiology of AVMs is not clear. Both congenital and posttraumatic forms have been observed. Although many AVMs are isolated, sporadic findings, they form an important aspect of some vascular malformation syndromes. A review of these syndromes and the underlying genetics has shed some light on mechanisms that can cause AVMs.

AVM: Hereditary Hemorrhagic Telangiectasia

Hereditary hemorrhagic telangiectasia (HHT) is a rare inherited vascular disorder affecting multiple vascular beds and associated with significant morbidity (Guttmacher et al. 1995; Govani and Shovlin 2009). This syndrome was originally differentiated from hemophilia by Henri Rendu in 1896 (Rendu 1896) and was well described in early case reports by Sir William Osler (Osler 1900) and Frederick Parkes Weber (Weber 1907) in the early 1900s. In recognition of these early descriptions, the condition is sometimes referred to as Osler-Weber-Rendu syndrome. HHT is characterized by spontaneous epistaxis, mucocutaneous telangiectasias, and visceral AVMs associated with a family history of the syndrome. These four components constitute the Curacao diagnostic criteria (Shovlin et al. 2000) for the condition. The presence of three or four of these components is diagnostic of HHT, whereas with one or fewer criteria the diagnosis is unlikely. Patients with HHT almost universally develop telangiectasias of the nasal mucosa and skin, which consist of dilated postcapillary venules with increased smooth muscle investiture often connected directly to dilated arterioles (Guttmacher et al. 1995). The genetic basis for HHT has been investigated, and mutations in any of three genes have been found to be associated with the condition. The majority of cases can be attributed to either endoglin (McAllister et al. 1994) (HHT type 1) or Activin receptor-like kinase 1 (Johnson et al. 1996) (ACVRL1—HHT type 2). A minority of cases is associated with mutations in SMAD4 (Gallione et al. 2004), in which there is clinical overlap with juvenile polyposis. All three of these genes are notable for their recognized roles in binding or signaling associated with the transforming growth factor-β (TGF-β) family of ligands (Govani and Shovlin 2009). Clinically significant AVMs of the brain, lung, liver, and GI tract are found in a significant minority of HHT patients (Guttmacher et al. 1995; Govani and Shovlin 2009).

The mutations that result in HHT include deletions, insertions, missense, nonsense, and splice-site mutations (Govani and Shovlin 2009) and are generally thought to result in loss of protein function. As more is known about the function of these proteins, mechanisms of AVM pathogenesis are being explored. Endoglin is an endothelial-specific cell surface protein (also known as CD105) that binds TGF-β ligands (McAllister et al. 1994) but has no known signaling functions. Matrix or cell-surface localization of these growth factors is a well-recognized mechanism essential for proper signaling, and loss of localizing proteins not only leads to HHT but is felt to be an important mechanism in Marfan syndrome (Neptune et al. 2003) as well. After the ligand is localized to the endothelial cell, signaling of TGF-β ligands involves binding to type II receptors (such as ACVRL1), which associate with and
phosphorylate type I receptors (Govani and Shovlin 2009). Upon phosphorylation, type II receptors become activated and phosphorylate downstream targets including the receptor-associated SMADs (SMADs 1, 2, 3, 5, and 8), which, in turn, bind with SMAD4 and translocate to the nucleus to affect gene transcription (Govani and Shovlin 2009). That the loss of endoglin or ACVRL1 results in a very similar clinical syndrome suggests that both genes lie in a common genetic pathway. Recent studies suggest that the probable ligands for this pathway are the bone morphogenetic proteins (BMPs) 9 and 10 (David et al. 2007; Scharpfenecker et al. 2007). Because heterozygosity for either endoglin or ACVRL1 is associated with HHT, it suggests that the disease results primarily from a decrease in signal from this common HHT pathway, as opposed to a new, pathologic signal from an alternative pathway.

Animal models have been developed to explore the in vivo functions of these proteins. The complete loss of either endoglin or ACVRL1 results in embryonic death in mid-gestation, with profound vascular abnormalities (Bourdeau et al. 1999; Li et al. 1999; Arthur et al. 2000; Oh et al. 2000; Urness et al. 2000). Interestingly, mice lacking either protein develop abnormal arteriovenous connections and shunts between the developing aorta and cardinal vein, and die as a consequence of inefficient circulation due to AVMs (Urness et al. 2000; Sorensen et al. 2003). Unlike humans, mice heterozygous for HHT proteins do not develop epistaxis or AVMs with predictably high penetrance. Indeed, it is common that mice with the same genetic lesion as humans do not share identical phenotypes. Although certain background mouse strains can encourage some features of HHT (Bourdeau et al. 1999), the disease is more profoundly manifested by the induced complete knockout of Acvr1l1 after critical stages of vascular development are allowed to proceed (Park et al. 2009). Using a tamoxifen-responsive Cre recombinase expressed in all tissues, including endothelium of all vascular beds, Park et al. (2009) recently showed the real-time development of AVMs in their mouse conditional mutant for Acvr1l1.

Although some vascular beds were prone to spontaneously develop AVMs, telangiectasias and AVMs of the skin only developed as a response to injury. This pattern has some analogy to HHT clinically, because AVMs in HHT patients are observed early and may be congenital in some vascular beds, such as brain, yet telangiectasias develop over time with predisposition for areas subject to trauma (Govani and Shovlin 2009).

AVM: Capillary Malformation–Arteriovenous Malformation

Recently another hereditary clinical vascular malformation syndrome has been described (Table 1) (Eerola et al. 2003) in which affected patients develop capillary malformations of the skin in a variety of locations, often associated with underlying AVMs or AV fistulae. The underlying high-flow lesions can occasionally alter the appearance of the otherwise erythematous capillary malformations, resulting in a pale halo surrounding the lesion (Boon et al. 2005). Some affected patients have more extensive involvement including hypertrophy and overgrowth of an affected limb (Parkes Weber syndrome). In light of these combined features, the syndrome has been termed CM-AVM. Families with CM-AVM have been found to have mutations in RASA1 encoding p120Ras-GAP (Eerola et al. 2003), a protein that regulates the activity of Ras.

Unlike HHT, in which the underlying mutations suggest abnormal growth factor signaling, the mutations in CM-AVM suggest a problem with signaling from small GTPases. Small GTPases are molecular switches that are well positioned to act as regulators of endothelial cell–cell interactions, junctional proteins, and the cellular cytoskeleton. Ras is a GTPase known to be important in linking signals from receptor tyrosine kinases (RTKs) at the cell surface to mitogen-activated protein kinase (MAPK) pathways in the cell to regulate growth, differentiation, and proliferation (Karnoub and Weinberg 2008). The importance of Ras in vivo is confirmed by the identification of increased levels of activated Ras in cancer cells. Since the
Identification of Ras, more than 100 further GTPases have been identified and characterized in separate families of the Ras "superfamily" of GTPases based on structural and functional characteristics. The Rho (Ras homolog) family GTPases were originally identified on the basis of homology with Ras and have been shown to control the cellular cytoskeleton and influence cell–cell interactions in several cell types including endothelial cells (Etienne-Manneville and Hall 2002).

<table>
<thead>
<tr>
<th>Syndrome</th>
<th>Associated malformations</th>
<th>Gene</th>
<th>Mutations</th>
<th>Phenotype</th>
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<tr>
<td>Hereditary hemorrhagic telangiectasia (HHT)</td>
<td>Arteriovenous malformations, mucocutaneous telangiectasias</td>
<td>Endoglin (HHT type 1)</td>
<td>Loss of function</td>
<td>Epistaxis, skin telangiectasias, lung, brain, and liver AVMs</td>
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<tr>
<td></td>
<td></td>
<td>ACVRL1 (HHT type 2)</td>
<td>Loss of function</td>
<td>Similar to type 1, but lung AVM less common, liver AVM more common</td>
</tr>
<tr>
<td></td>
<td></td>
<td>SMAD4 (HHT/JP)</td>
<td>Loss of function</td>
<td>Overlap syndrome of typical HHT plus juvenile polyposis</td>
</tr>
<tr>
<td>Cerebral cavernous malformations (CCMs)</td>
<td>Cerebral and retinal cavernous malformations, cutaneous capillary malformations, and hyperkeratotic cutaneous capillary venous malformations</td>
<td>KRIT1 (CCM1)</td>
<td>Loss of function</td>
<td>Multiple brain CCMs, high disease penetrance by MRI screening, subset asymptomatic</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CCM2</td>
<td>Loss of function</td>
<td>Similar to CCM1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PDCD10 (CCM3)</td>
<td>Loss of function</td>
<td>More severe phenotype, earlier onset, smaller families</td>
</tr>
<tr>
<td>Capillary malformation–arteriovenous malformation (CM-AVM)</td>
<td>Capillary malformations, arteriovenous malformations, arteriovenous fistulae, Parkes Weber syndrome</td>
<td>RASA1</td>
<td>Loss of function</td>
<td>Capillary malformations, some with pale &quot;halo&quot; caused by vascular steal from underlying high-flow AVM. Rare family members have associated hypertrophy of an affected limb (Parkes Weber syndrome).</td>
</tr>
<tr>
<td>Cutaneomucosal venous malformations (VMCMs)</td>
<td>Multifocal mucocutaneous small venous lesions, other venous malformations</td>
<td>TEK</td>
<td>Missense, activating mutations</td>
<td>Variability from cosmetic vascular lesions to multifocal venous malformations affecting vital organs</td>
</tr>
</tbody>
</table>

These proteins are activated when bound to GTP and then become inactivated as they hydrolyze GTP to GDP. The dynamics of GTP binding and hydrolysis, as well as the subcellular localization of GTPases, contribute to the biologic effects of these systems. A variety of associated proteins modulate GTPase activity by promoting GDP–GTP exchange (guanine-nucleotide exchange factors, or GEFs) or by promoting GTP–GDP hydrolysis (GTPase-activating proteins, or GAPs). Scaffold proteins
contribute to GTPase regulation by bringing the GTPase switch into proximity with downstream signaling effectors (Jaffe et al. 2005), and post-translational isoprenylation allows GTPases to localize with membranes correctly. The potential complexity of GTPase signaling is staggering, and in vitro studies cannot confirm the importance of individual pathways in vivo. Human vascular malformation syndromes such as CM-AVM have highlighted GTPase-associated pathways that are of particular importance to vascular morphogenesis and stability in vivo.

The mutations found in CM-AVM result in a loss of function of p120-RasGAP, which is a GTPase-activating protein. Because GTPase activity leads to hydrolysis of GTP to GDP and inactivates signaling, the loss of p120-RasGAP opens the potential for increased Ras activation. RASA1 has homology with neurofibromin (NF1), the gene mutated in cases of neurofibromatosis, yet NF1 expression is not sufficient to compensate for loss of RASA1 in families with CM-AVM (Boon et al. 2005). The importance of RASA1 relative to NF1 or two other homologous genes, RASA2 and RASAL, in vascular stability would not have been predicted on the basis of in vitro studies and was clearly shown by human genetic studies.

In addition to its function as a Ras-GAP, p120-RasGAP has been shown to associate with p190-RhoGAP, where it may influence Rho kinases to modulate the cellular cytoskeleton (Settleman et al. 1992a,b). Furthermore, p120-RasGAP associates with Rap1a (Freh et al. 1990), a GTPase often associated with cellular adhesion and cell–cell interactions, and the bait used to initially identify KREV1/RAP1a interaction trapped 1 (KRIT1) (Serebriiskii et al. 1997), a gene subsequently found to be mutated in a subset of families with cerebral cavernous malformations (CCMs) (Laberge-le Couteulx et al. 1999; Sahoo et al. 1999).

Cerebral Cavernous Malformations

Cerebral cavernous malformations (CCMs) are common vascular malformations described predominantly in the central nervous system. Lesions consist of dilated endothelial channels or caverns lacking smooth muscle support and filled with blood or thrombus (Labauge et al. 2000). Occasionally, CCMs rupture, leading to hemorrhagic stroke or death. Even in the absence of overt hemorrhage, all lesions are associated with hemosiderin, a blood breakdown product in the surrounding brain parenchyma. Ultrastructural studies suggest absent or diminished tight junctions, implying localized loss of the blood–brain barrier (Clatterbuck et al. 2001).

CCM lesions are found with a prevalence of one in 200 people (Otten et al. 1989; Vernooij et al. 2007). Approximately 20% of patients have familial disease, following an autosomal-dominant pattern of inheritance. CCM has been linked to loss-of-function mutations in the genes encoding any of three structurally distinct proteins—KRIT1 (CCM1) (Laberge-le Couteulx et al. 1999; Sahoo et al. 1999), CCM2 (Osmosensing Scaffold for MEKK3, OSM; Malcavernin; or MGC4607) (Liquori et al. 2003; Denier et al. 2004), or Programmed cell death 10 (PDCD10; CCM3) (Bergametti et al. 2005). For a more complete review of CCM genetics, see Riant et al. (2009). The three CCM disease genes are structurally unrelated intracellular proteins that lack catalytic domains and have been found to associate with one another, and influence a variety of signaling pathways. This topic was recently reviewed in greater detail by Faurobert and Albiges-Rizo (2009).

Initial evidence that CCM might result from abnormal GTPase activity came with the identification of KRIT1 mutations in families with CCM and linkage to CCM1. Further proof that the KRIT1–RAP1a interaction is functionally important came from studies in endothelial cells in vitro (Glading et al. 2007) and Krit1 heterozygous knockout mice in vivo (Glading and Ginsberg 2010). KRIT1 and RAP1a interact to influence endothelial cell junctional integrity through downstream signaling cascades such as β-catenin signaling. Krit1 has also been shown to interact genetically with the related GTPase, Rap1b (Gore et al. 2008).

CCM2 was identified simultaneously as a gene associated with familial CCM and as a scaffold protein to facilitate stress signaling from the

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small GTPase Rac1 to p38 MAPK (Uhlik et al. 2003). CCM2 also associates with RhoA, and the loss of CCM2 leads to increased RhoA activity in the endothelium (Whitehead et al. 2009). Increased RhoA activity is associated with vascular instability, and favorable effects of RhoA inhibition in CCM2 deficiency are seen both in vitro and in vivo, suggesting a possible therapeutic strategy for CCM disease (Whitehead et al. 2009). A similar mechanism appears to be present in KRIT1 deficiency (Stockton et al. 2010).

Until recently little was known about PDCD10. This gene was initially identified in a screen for proteins associated with apoptosis. As with KRIT1 and OSM, until the protein was linked to CCM disease, there was little to suggest the importance of this gene in the vasculature by structure, homology, or expression pattern (Petit et al. 2006). Overexpression studies have found that PDCD10 can be found in association with KRIT1 and OSM (Hilder et al. 2007), suggesting a possible common mechanism. Increased RhoA activity has also been observed with the loss of PDCD10 (Zheng et al. 2010). However, rather than binding to GTPases and MAPKs, PDCD10 has been found to associate with serine-threonine kinases (STKs), including STK24, STK25, and MST4, and a further complex of associated proteins (Goudreault et al. 2009). Whether this complex represents a unique, PDCD10-associated signaling mechanism, or whether with endogenous levels of protein there is cross talk between PDCD10 and KRIT1, OSM and GTPase pathways is not known. The loss of any of these three proteins results in a common vascular pathology, yet it is not clear whether CCM represents the end stage of a single pathogenic process or an unexpectedly common result of distinctly different (genotype-specific) pathways.

The GTPase pathways associated with CM-AVM and CCM regulate vascular permeability and control endothelial cell–cell interactions. With this observation, a theme is introduced that the loss of proteins that regulate vascular stability results in vascular malformations. This is particularly well illustrated in the case of cutaneomucosal venous malformations.

Cutaneomucosal Venous Malformations (VMCMs)

Venous malformations (VMs) are the most common of vascular malformations and account for >50% of individuals reporting to vascular malformation centers. VMs are the result of local aberrations in angiogenesis occurring in fetal development and can range in clinical spectrum from benign cosmetic variocities to multifocal lesions in multiple vital organs (Brouillard and Vikkula 2003; Wouters et al. 2009). The majority of vascular malformations have no known genetic component and are sporadic in nature. However, families with dominant inheritance of VMs have been identified, and linkage analysis has localized mutations in the TEK gene on chromosome 9p21. This gene codes for the angiopoietin-1 receptor protein Tie2, and the majority of mutations identified thus far have been in or near the amino-terminal-most of two kinase domains within the intracellular portion of the protein (Wouters et al. 2009). These autosomal-dominant venous malformations termed cutaneomucosal venous malformations (VMCMs) tend to involve small multifocal mucocutaneous lesions in addition to the possibility of other venous malformations. Although they only make up ~2% of reported venous malformations, the understanding of the genetic factors leading to VMCMs have led to partial understanding of the much more common sporadic venous malformations (Limaye et al. 2009a).

To date, 17 families have been reported with mutations in or near the kinase domain of Tie2, and roughly 60% of these are missense mutations causing an arginine–tryptophan substitution (R849W) (Limaye et al. 2009b). When exogenously expressed in human umbilical vein endothelial cells (HUVECs), this mutation had no effect on migration or cell proliferation when compared with its wild-type counterpart. However, Tie2-R849W-expressing cells were significantly more resistant to apoptosis by serum withdrawal and formed unstable tubes (Hu et al. 2008), suggesting a possible mechanism for the propagation of vascular malformations that possess a paucity of smooth muscle or
pericyte support that would normally induce endothelial apoptosis (Benjamin et al. 1999). Additionally, Hu et al. (2008) determined that Tie2-R849W was hyper-autophosphorylated and, unlike its wild-type counterpart, activated the signal transducer and activator of transcription 1 (STAT1) transcription factor in an Ang1-independent but Janus kinase (JAK)–dependent manner. Intriguingly, however, STAT1 activation has actually been shown to down-regulate angiogenesis in vivo (Battle et al. 2006), suggesting that further evaluation of STAT1 signaling in VMCM patients is warranted and this pathway could be a potential therapeutic target in treating vascular malformations.

Another insight into the understanding of venous malformations has come from the discovery of a “second hit” within the VMCM of an individual with the inherited R849W mutation (Limaye et al. 2009b). Under the hypothesis of a paradigmatic mode of inheritance for VMCM, they found a Tie2 deletion mutation in the vascular lesion of a single patient that did not correspond to the genotype of DNA isolated from the blood of this patient. This finding prompted the investigators to sequence DNA from vascular malformations of other patients whose VMs were inconsistent with the dominant inherited VMCM. In screening 62 VMs from 57 patients for mutations in TEK (Tie2), the investigators found 30 mutations in 28 individuals. Upon further characterization of the most prominent of these mutations, a leucine–phenylalanine substitution (L914F) within the same kinase domain as the inherited R849W mutation, they determined that this mutation also conferred hyper-autophosphorylation of the Tie2 receptor, suggesting a similar mechanism for the development of sporadic VMs as to that of VMCMs (Limaye et al. 2009b).

CONCLUDING REMARKS

The vascular system is remarkably well regulated: able to remain stable in established tissues, yet immediately responsive and able to sprout and migrate to perfuse new tissues that form in response to injury or growth. Vascular morphogenesis is normally well controlled, and vascular malformations are fortunately rare. The important role of pro-angiogenic factors such as VEGFs and FGFs in vascular biology has long been appreciated. Vascular malformation syndromes as discussed in this article highlight an underappreciated mechanism in the control of vascular biology—that of stabilizing factors that counteract pro-angiogenic signals. That such mechanisms exist is not surprising. It is a recurring theme in nature that opposing signals are involved in regulating biologic processes, and the balance between opposing forces determines the net effect. Just as the parasympathetic nervous system is the ying to the sympathetic nervous system yang, or that pro-thrombotic pathways are generally held in check by anti-thrombotic mechanisms, thus do pathways of vascular stability predominate in a stable vessel to prevent pathologic vascular leak and excessive angiogenesis and neovascularization. As more stabilizing pathways are appreciated and described, the key proteins involved become candidate genes for these and other vascular malformation syndromes. Similarly, there is a strong possibility that genes associated with vascular malformations are involved in regulating vascular stability.

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