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Intracranial aneurysms: links among inflammation, hemodynamics and vascular remodeling

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Abnormal vascular remodeling mediated by inflammatory cells has been identified as a key pathologic component of various vascular diseases, including abdominal aortic aneurysms, brain arteriovenous malformations and atherosclerosis. Based on findings from observational studies that analysed human intracranial aneurysms and experimental studies that utilized animal models, an emerging concept suggests that a key component of the pathophysiology of intracranial aneurysms is sustained abnormal vascular remodeling coupled with inflammation. This concept may provide a new treatment strategy to utilize agents to inhibit inflammation or cytokines produced by inflammatory cells such as matrix metalloproteinases. Such an approach would aim to stabilize these vascular lesions and prevent future expansion or rupture. [Neurology 2006; 28: 372–380]

Keywords: Intracranial aneurysms; inflammation; remodeling; pathophysiology

Approximately 27,000 Americans are estimated to suffer subarachnoid hemorrhage resulted from ruptured intracranial aneurysms each year. Ruptured intracranial aneurysms often result in catastrophic consequences causing severe morbidity and high mortality. Despite recent advancements in the diagnosis and treatment of intracranial aneurysms, mechanisms for the formation, development and subsequent rupture of intracranial aneurysms are not well understood.

Abnormal vascular remodeling mediated by inflammatory cells has been identified as a key pathologic component of various vascular diseases, including abdominal aortic aneurysms, brain arteriovenous malformations (BAVM) and atherosclerosis. This concept may provide a new treatment strategy to utilize agents to inhibit inflammation or cytokines produced by inflammatory cells such as matrix metalloproteinases (MMPs). Such an approach would aim to stabilize these vascular lesions and prevent future expansion or rupture (Figure 1).

Based on findings from observational studies that analysed human intracranial aneurysms and experimental studies that utilized animal models, an emerging concept suggests that a key component of the pathophysiology of intracranial aneurysms is sustained abnormal vascular remodeling coupled with inflammation.

HEMODYNAMIC STRESS AND INTRACRANIAL ANEURYSMS

Vascular remodeling is intimately tied to hemodynamics. Blood vessel is an active organ that can undergo a dynamic remodeling process in response to acute and chronic changes in blood flow, blood pressure and metabolism. Sustained hemodynamic stresses, especially elevated wall shear stress at the bifurcation or the outer lateral curve of blood vessels, may trigger physiologic vascular remodeling as an adaptive process.

Intracranial aneurysms are commonly found in the location where abnormal hemodynamic stresses are exerted on the vascular wall. These locations include the junction of the internal carotid artery and the posterior communicating artery, the anterior communicating artery complex and the bifurcation of the basilar artery. Vascular remodeling triggered by abnormal hemodynamic stress on the blood vessels inside the circle of Willis may play a role in the formation, development and subsequent rupture of intracranial aneurysms. Increased hemodynamic stresses especially elevated wall shear stress at the bifurcation or the outer lateral curve of blood vessels may trigger physiologic vascular remodeling as an adaptive process.
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A number of studies have estimated the hemodynamic stresses on the cerebrovasculature and aneurysm wall by analysing either idealized aneurysm models or clinical data obtained by angiography or magnetic resonance imaging technique\(^\text{15-19}\). Various mathematical models were used to estimate the wall shear stress and pressure. These studies indeed predicted that an apex of bifurcation or the outer lateral wall of a curved blood vessel, common sites of intracranial aneurysms, have abnormally high hemodynamic stresses, i.e. high pressure and shear stress. Such hemodynamic stresses may lead to the initiation of normal adaptive outward vascular remodeling.

However, owing to complex geometry of these locations, adaptive flow-induced outward vascular remodeling becomes asymmetric remodeling that can be recognized as a pre-aneurysm change. Once such pre-aneurysm starts forming, the distal neck of aneurysm wall will have sustained hemodynamic stress\(^\text{15,17}\). Asymmetric and sustained outward vascular remodeling will eventually lead to the growth of aneurysms.

The ‘impact zone’, the location on the aneurysmal wall exposed to high hemodynamic stresses, can be better characterized by elevated local wall shear stress than elevated local pressure\(^\text{15}\). The flow impact results in two orthogonal forces. The direct ‘impacting force’ or ‘dynamic pressure’ at the stagnation point is perpendicular to the vessel wall and the wall shear stress, the viscous friction from the bloodstream that passes next to the stagnation point, is tangential to the wall (Figure 2).

Although it has been suggested that the ‘impacting force’ or the resulting local pressure elevation may contribute significantly to aneurysm development\(^\text{20-23}\), some studies suggested that such focal pressure elevations are rather insignificant or may not be a critical physical force. Two studies\(^\text{24,25}\) reported that the local pressure elevations at branch points and bends in arteries were \(~1\) mmHg, which is only 1–2% of the peak intravascular pressure. The pressure elevation inside the aneurysmal sac produced by flow impact was found to be even weaker\(^\text{24}\). However, substantial wall shear stresses were found beside the direct impingement point; 200 dynes/cm\(^2\) at the apex of an arterial bifurcation\(^\text{25}\) and 100 dynes/cm\(^2\) at the distal neck of a saccular aneurysm\(^\text{15}\). Consistent with the common notion that the upstream triggers for flow-mediated vascular remodeling events come from endothelial cells, sensors for wall shear stress\(^\text{26-30}\), Hoi et al. define the impact zone as the vascular area that experiences a wall shear stress above the physiological level of 20 dynes/cm\(^2\) (Ref. \text{15}).

Another factor that sometimes is classified as a hemodynamic stress is systemic or essential hypertension, which is not the local pressure elevation owing to local hemodynamic conditions that was previously discussed. As will be discussed as either a cause or effect of the disease, systemic hypertension represents a systemic endothelial dysfunction, in addition to the purely hemodynamic effects on the vascular tree.

Systemic hypertension is a consistent risk factor for intracranial aneurysm development\(^\text{31,32}\). Like other risk factors such as smoking and genetic disposition\(^\text{1}\), it is not well understood how systemic hypertension contributes to aneurysm development. From a very basic mechanical point of view, intravascular pressure subjects the vessel wall under tensile stress\(^\text{33}\), which could cause the aneurysmal sac to enlarge if the wall is already degraded.

However, this simple view does not uniquely explain how systemic hypertension contributes to the vessel wall degradation that is essential for aneurysm initiation, growth and rupture.

Research on systemic hypertension and peripheral arteries provides some clues as to how systemic hypertension may affect the arterial wall functions.

Hypertension appears to be associated with impairment of endothelial function in its vasodilation response
to acetylcholine infused into the brachial artery\textsuperscript{34,35}. For example, two studies showed that endothelium-dependent vasodilation by acetylcholine is reduced in the forearm of essential and secondary hypertensive patients\textsuperscript{34,35}. An altered balance of vasodilator and vasoconstrictor molecules could probably characterize hypertension-related endothelial dysfunction.

In patients with hypertension, the endothelial cell vasodilatory response that includes prostacyclin and nitric oxide production was diminished\textsuperscript{34}, while the production of endothelin-1\textsuperscript{36}, reactive oxygen species, angiotensin II\textsuperscript{37} and other endothelium-derived constricting factors was either maintained or increased. It has also been found that chronic arterial hypertension results in altered intercellular adhesion molecule-1 (ICAM-1) expression on the endothelium, which may contribute to the abnormal inflammatory responses associated with this disease\textsuperscript{38}. The altered pathologic behavior of the vasculature may contribute to intracranial aneurysm formation and development by preventing a healthy remodeling process in response to a sudden increase of blood flow.

High shear stress and hypertension alone may not be sufficient to explain the hemodynamic conditions responsible for intracranial aneurysm development. It is well established that normal vascular remodeling is an arterial adaptation to a long-term increase or decrease in wall shear stress from the baseline level of 15–20 dynes/cm\textsuperscript{2} (Ref. 39). Increased flow causes dilation of the artery until the wall shear stress is restored to baseline\textsuperscript{11,40}. The remodeling process of the common carotid artery in response to high flow and shear stress is well described in the work of Sho et al.\textsuperscript{41}. An increase in the early expression of MMP-2 and -9 is associated with a sudden increase in blood flow and wall shear stress\textsuperscript{41}. Inhibition of MMPs by doxycycline was able to reduce outward remodeling in response to an increased blood flow.\textsuperscript{12} The resulting degradation of internal elastic lamina allows the artery to be dilated and the wall shear stress to be reduced, followed by the proliferation of endothelial cells and smooth muscle cells\textsuperscript{41}. This remodeling mechanism does not seem to lead to a diseased condition. Other than the degraded internal elastic lamina, the remodeled wall of the arterial tube was rather uniform and healthy with abundant endothelial and smooth muscle cells.

The morphology of the saccular intracranial aneurysm wall, nonetheless, is quite different. Besides having a disrupted internal elastic lamina, the aneurysm wall is very inhomogeneous; endothelial layer is usually disrupted; media is thinned and smooth muscle cells are scarce, evidently owing to apoptosis and possibly by inhibition of proliferation; inflammatory cells are frequently found\textsuperscript{42}. Such remodeling appears to be related to the complex hemodynamic microenvironment associated with the apex or flow divider. This particular geometry harbors a ‘disturbed flow’ created by flow impingement in addition to the high shear stress (Figure 3). The high shear stress might play the critical role in degrading internal elastic lamina through the release of MMPs, but the ‘disturbed flow’ may be just as important for aneurysm development as the high shear stress.

The ‘disturbed flow’ has been a key concept in the onset of atherosclerosis, stenosis, thrombosis and some inflammation\textsuperscript{18,39,43}. In these situations, ‘disturbed flow’ is generally used to describe a recirculating flow in a cavity or wall expansion near an arterial branch. The shear stress is low owing to the slow velocity, but the change in direction is frequent\textsuperscript{44}. Blood elements stay in the region for a prolonged period, greatly increasing the adhesion of elements such as platelets and monocytes to endothelial cells\textsuperscript{43}. It should be noted that near the reattachment point of the recirculation bubble, the flow splits into opposite directions; therefore, the local shear stress gradient (spatial variation of shear stress) is extremely high. Increasing evidence suggests that it is not the low shear stress but the high wall shear stress gradient that triggers the modification of endothelial gene expression, causing the cascade of events that lead to atherosclerosis, stenosis and related inflammation\textsuperscript{15–47}.

The hemodynamic environment associated with saccular aneurysms is also likely to constitute a ‘disturbed flow’, whereby the flow splits into opposite directions near the impingement point and produces a high spatial variation of shear stress or wall shear stress gradient. This ‘disturbed flow’ may distinguish the pathologic remodeling that leads to saccular aneurysm development from the normal outward remodeling of an arterial tube subjected to increased flow. It is possible that, together with the high shear stress and high pressure, high wall shear stress gradient mediates saccular aneurysm formation.
Fusiform aneurysms constitute ~2% of all intracranial aneurysms. Unless the aneurysmal distention is severely asymmetric (a case bordering with the saccular aneurysm), no impinging flow is usually present and hemodynamic stress is usually low in a fusiform aneurysm. We nevertheless find the ‘disturbed flow’—the recirculation flow in the expansion area. Such recirculation flow is similar to that in the abdominal aortic aneurysm and atherosclerosis development\(^\text{19}\). Indeed, fusiform aneurysms are known as atherosclerotic aneurysms\(^\text{18,49}\).

In order to further characterize exact relationships between hemodynamic stresses and a clinical course of intracranial aneurysms, a means to measure shear stress and pressure in the human cerebral circulation needs to be further developed\(^\text{19,24}\).

**ANIMAL MODEL OF INTRACRANIAL ANEURYSM INDUCED BY HEMODYNAMICS**

A critical role of hemodynamic stresses in the formation of intracranial aneurysms has been demonstrated by studies using an animal model of intracranial aneurysms. Experimentally, changing hemodynamic stresses in the circle of Willis can induce intracranial aneurysms. In animals, a combination of unilateral carotid ligation and hypertension can cause intracranial aneurysms in the circle of Willis through an increase in blood flow and local arterial pressure. This model was originally developed and has been extensively studied by a group of researchers led by Dr. Nobuo Hashimoto at Department of Neurosurgery at Kyoto University in Japan. In this animal model, aneurysms were frequently discovered in areas of high hemodynamic stresses such as at (1) the bifurcation between anterior cerebral artery and olfactory artery contralateral to the ligated common carotid artery and (2) posterior communicating arteries\(^\text{50}\).

Interestingly, rats that underwent bilateral common carotid ligation displayed higher incidences of intracranial aneurysms in the posterior circulation than those that underwent unilateral ligation\(^\text{50}\). Presumably, rats that underwent bilateral common carotid artery ligation developed higher blood flow through the basilar artery to posterior communicating arteries than those with unilateral carotid ligation, because a majority of the blood flow to the brain needs to be supplied by basilar artery in the rats that underwent bilateral ligation. This finding may indicate a dose-relationship between an increase in the blood flow and the incidence of aneurysms.

Histologically, these hemodynamically-induced intracranial aneurysms in rats closely resemble intracranial aneurysms in humans. Elastic lamina in this model showed degenerative changes such as loss or fragmentation of the lamina, which are common in human intracranial aneurysms\(^\text{15,51}\). In addition, both human intracranial aneurysms and rodent experimentally-induced aneurysms showed damage to endothelial layers and thrombi in the sac\(^\text{52}\). Death of medial smooth muscle cells through apoptosis was also observed in the rat aneurysms\(^\text{53}\).

The same manipulation of intracranial hemodynamics in rats resulted in intracranial aneurysm formation in mice and primates\(^\text{54-57}\). It should be noted that some studies using these models utilized beta-aminopropionitrile (BAPN) treatment to increase the incidence of aneurysms. BAPN is an irreversible inhibitor of lysyl oxidase that inhibits the crosslinking of collagen and elastin fibers, which are critical structural components of blood vessels. Some histologic changes in the structural components of the vascular wall, such as the disruption or loss of elastic lamina in these models, may be due to the inhibition of the normal metabolism of collagen and elastin fibers by BAPN treatment.

The addition of BAPN treatment to the combination of unilateral carotid ligation and hypertension had quadrupled the incidence of aneurysms\(^\text{58}\). This may be analogous to the increased incidence of aneurysms in patients with Ehlers-Danlos syndrome, in which common defects are in collagen synthesis and metabolism.

**INFLAMMATION IN HUMAN INTRACRANIAL ANEURYSMS**

The potential roles of inflammation in the development and rupture of intracranial aneurysms have been suggested by observational studies that analysed the presence of inflammatory cells or markers in human intracranial aneurysm tissues or serum samples from patients harboring intracranial aneurysms. Several studies reported the presence of inflammatory cells in the wall of intracranial aneurysms. Histologic analysis of intracranial aneurysms using electron microscopy showed the presence of extracellular lysosome-like granules and leukocytes near disorganized elastic lamina, indicating degradation of elastic lamina by lytic enzymes secreted by leukocytes in the vascular wall\(^\text{59}\).

Analysing mostly unruptured aneurysm tissues (23 unruptured and two ruptured aneurysms), Chyatte et al. reported the presence of macrophages/monocytes and T-lymphocytes in the aneurysmal wall\(^\text{7}\). Previously, a similar observation was attributed to potential atherosclerotic changes in part of the aneurysmal sac\(^\text{60}\). However, Chyatte et al. found these inflammatory cells scattered throughout the aneurysmal wall, indicating that inflammation in intracranial aneurysms is not limited to the part of the vascular wall undergoing atherosclerotic changes but rather is a generalized reaction. Kataoka et al. examined leukocyte infiltration in relation to structural changes of the aneurysmal wall in both ruptured and unruptured intracranial aneurysms\(^\text{32}\). Half of the unruptured aneurysms (10/20) and all of the ruptured aneurysms (40/40) had been infiltrated by leukocytes. Leukocyte infiltration appeared to be associated with the damage or loss of the smooth muscle cell layer and collagen fibers.

Similar findings were reported by Frösen et al.\(^\text{8}\). In this study, 24 unruptured and 42 ruptured aneurysms were analysed. The majority of both unruptured and ruptured aneurysms showed evidence of macrophage and leukocyte infiltration, with more prominent leukocyte infiltrations in the aneurysms that had ruptured.
Interestingly, in unruptured aneurysms, the areas on the vascular wall that showed macrophage and leukocyte infiltration were also free of organizing thrombosis or myointimal hyperplasia, changes often regarded as atherosclerotic. This indicates that inflammation mediated by these inflammatory cells in the aneurysmal wall is not a necessary part of atherosclerosis or the tissue repairing process after rupture.

The associations in these studies between histologic findings and rupture need to be carefully interpreted. The cross-sectional nature of these studies makes the exploration of a causal relationship between inflammation and aneurysmal rupture complex. Part of the challenge is distinguishing if structural changes in the vascular wall and leukocyte infiltration in aneurysms are due to the rupture itself or an epiphenomenon. For example, the inflammation and small degree of remodeling that follows an aneurysmal rupture could be due to the provocation of intravascular thrombosis and normal tissue repair. However, this does not explain the presence of inflammatory cells in unruptured aneurysms and as several studies have consistently found it, the majority of both ruptured and unruptured intracranial aneurysms exhibited macrophage and lymphocyte infiltration into the vascular wall.

The presence of inflammation in intracranial aneurysms is further supported by studies that analyzed cytokines related to inflammation in serum samples from patients with intracranial aneurysms. Serum elastase and collagenase levels seem to be elevated in patients with intracranial aneurysms. These findings were corroborated by studies intracranial aneurysm tissues for expression or activity of collagenase and elastase. High levels of collagenases and elastases were observed in intracranial aneurysms tissues. The major collagenases in intracranial aneurysm tissues seem to be MMP-2 and -9 (Ref. 67). MMP-2 and -9 are known to be produced by inflammatory cells, especially by macrophages in other vascular diseases such as abdominal aortic aneurysms.

Predicting natural course of aneurysms in regard to a propensity to rupture using serum levels of these serum proteinases may be intriguing, but data for associations between rupture and the serum proteinase levels are conflicting. To explore the relationship between the presence of serum proteinases and aneurysmal rupture, a longitudinal study following patients with unruptured aneurysms is needed.

**HEMODYNAMIC STRESS AND INFLAMMATION**

Hemodynamic stress can trigger an inflammatory process by activating endothelial and inflammatory cells. High shear stress, i.e. high blood flow, activates endothelial cells and up-regulates leukocyte adhesion molecules including ICAM-1 and chemokines such as monocyte chemotactic protein-1 (Refs. 69–72). These molecules attract circulating neutrophils and monocytes, which facilitate their invasion into the vascular wall and their maturation. Along with activated endothelial cells and smooth muscle cells, these inflammatory cells secrete cytokines, including MMPs and elastases, and initiate outward vascular remodeling.

Other studies have linked the ‘disturbed flow’ pattern to the increased leukocyte adhesion to endothelial cells and ultimately to the observed inflammation and atherosclerotic lesions in the arterial wall. The flow disturbance may cause an increased residence time of circulating particles in the region and increased mass transfer, therefore enhancing the interactions between circulating leukocytes and vascular endothelial cells. Furthermore, there is considerable evidence that spatial variations of shear stress in the ‘disturbed flow’ region exert significant influences on endothelial cells. The adhesion of monocytes on endothelial cells occurs preferentially in the vicinity of the reattachment point in a recirculating flow. The large shear stress gradient not only increases monocyte adhesion to endothelial cells but also causes a net migration of endothelial cells away from the region of high shear gradient.

We have pointed out that both saccular and fusiform aneurysms experience a ‘disturbed flow’. The flow splitting at the impingement point of the inflow jet or the reattachment point of recirculating flow creates large gradients in fluid shear stress on the luminal surface of the aneurysm, the neck and possibly the immediately adjacent parent vessel. Such large shear gradients may be responsible for monocyte adhesion. In addition, according to the study of Munn et al., a flow velocity component perpendicular to the vascular wall can cause a cell flux and therefore lead to direct invasion of large blood elements into the wall. Flow impingement in saccular aneurysms provides such perpendicular velocity to the wall, which, coupled with the possible endothelial migration away from this region of large spatial gradient of shear stress, may further facilitate leukocyte invasion.

Furthermore, both saccular and fusiform aneurysms contain recirculating flow inside their cavities, where the long residence time of particles enhances the mass transfer between the circulating particles and the vascular endothelial layer. The recirculation zone (cavity) is therefore the pro-atherosclerotic area.

**HEMODYNAMICS, VASCULAR REMODELING, INFLAMMATION AND THE DEVELOPMENT OF INTRACRANIAL ANEURYSMS**

Based on experimental and observational findings previously discussed, a concept that views the formation of intracranial aneurysms as a sustained and focalized vascular remodeling emerges (Figure 4). While normal outward vascular remodeling in response to an increase in blood flow is a healthy adaptive process to normalize the wall shear stress, aneurysm development is a pathologic manifestation of failure to re-establish homeostasis under hemodynamic insults.

Inflammation has been found to play an important role in this process, whether by contributing to the initiation of the cascade for the outpouching process,
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Figure 4: Schematic representation of normal outward vascular remodeling leading to aneurysm formation at a flow divider. Increased hemodynamic stresses activate endothelial cells, which cause recruitment and activation of inflammatory cells and secretion of MMPs to degrade the internal elastic lamina. These processes initiate outward remodeling. In normal vascular remodeling, the luminal diameter increases. The vessel enlargement brings the shear stress down to baseline level; therefore, blood vessels will be adapted and stabilized. However, during aneurysm formation, outward vascular remodeling becomes asymmetric or focalized. Flow impingement creates a complex hemodynamic environment with spatially varying shear stress and increased invasion of inflammatory cells. As the micro-aneurysm grows, part of the wall will continue to experience high shear stress. Focalized vascular remodeling continues with increasing participation of inflammation, which could repair the wall, stabilize the aneurysm or further degrade it, leading to rupture attempting to repair the wound or accelerating the degradation of the wall en route to aneurysm rupture.

Intracranial aneurysm formation may start as an otherwise normal outward vascular remodeling at a bifurcation point or curve. However, because of the particular geometry of the affected blood vessels (branching point or curved blood vessel), outward vascular remodeling during the development of intracranial aneurysms may become asymmetric or focalized. The focus of the vascular remodeling causes pre-aneurysmal change and leads to microaneurysm formation, a small bulging of the blood vessel. As the micro-aneurysm grows, part of it will be subjected to sustained high shear stress. In this area, endothelial cells are kept either activated or damaged, causing the accumulation of inflammatory cells. Focalized vascular remodeling continues with increasing inflammation. When the inflammation is intense, major structural components of the vascular wall may be destroyed and lead to aneurysm rupture. A hallmark of the pathophysiology of intracranial aneurysms may be a sustained, focalized outward vascular remodeling coupled with inflammation.

Genetic factors, such as certain single nucleotide polymorphisms or other underlying diseases including hypertension and atherosclerosis, may have a role in accentuating the sustained vascular remodeling and inflammation.

RELATED CEREBROVASCULAR CONDITIONS: BAVM
There may be a parallel in the inflammation observed in the walls of aneurysms operative in patients harboring BAVM. Lacking intervening capillaries, BAVM function as arteriovenous shunts in the cerebro-circulation. Abnormal blood vessels in BAVM (nidal vessels) are exposed to high blood flow and possibly high shear stress. Although their genesis is not well understood, there is growing evidence that suggests a role for sustained vascular remodeling and inflammation in the pathophysiology of BAVM.

Genetic and tissue analysis in BAVM patients show an interesting link between inflammatory cytokines and the clinical behavior of BAVM. Interleukin-1 (IL-1) and tumor necrosis factor-alpha (TNF-α) are potent inflammatory cytokines that can serve as tumor and endothelial cell growth factors. The GG genotype of the IL-6-174C promoter polymorphism is associated with clinical presentation of intracranial hemorrhage (ICH) in BAVM patients. Further, TNF-α-238G>A promoter polymorphism is associated with an increased risk of new hemorrhage in the natural course of BAVM patients.

Chen et al. investigated whether tissue IL-6 expression was associated with IL-6-174C>G genotype and IL-6 was linked to putative downstream targets involved in angiogenesis and vascular instability. They found that the highest IL-6 protein levels in BAVM tissue were associated with IL-6-174GG genotype. IL-6 induced MMP-3 and -9 expression and activity in normal mouse brain and increased proliferation and migration of cerebral endothelial cells. Together, these results suggest that the IL-6 genotype associated with ICH appears to have functional differences in IL-6 tissue expression, consistent with the hypothesis that inflammatory processes induce angiogenic activity and possibly contribute to BAVM ICH.

Because BAVM tissue expresses both MMP-9 (Ref. 3) and IL-6 (Ref. 85), the question arises as to from which cell population those cytokines were primarily derived. Preliminary evidence suggests that it is by infiltrating leukocytes, in particular neutrophils.

Taken together, it appears that a combination of genetic and local environmental factors helps shape the
BAVM lesional phenotype and appears to be responsible for some portions of the clinical behavior. More studies need to be conducted to dissect their operative mechanisms as well as their relationship to the actual pathogenesis of the disease. In any event, there may be a broad overlap in the inflammatory contribution to the phenotype of both aneurysms and BAVM.

FUTURE DIRECTIONS
Findings from observational and experimental studies suggest roles for vascular remodeling and inflammation in the pathophysiology of intracranial aneurysms. However, there is a lack of study linking these two processes, i.e. vascular remodeling and inflammation in this pathology. A recent advancement in image acquisition and analysis techniques that enable a longitudinal follow-up of changes in geometry of intracranial aneurysms and local hemodynamic stresses in humans will help us to study the effects of changing hemodynamic stresses on aneurysmal growth and subsequent rupture. Animal models, especially hemodynamically-induced intracranial aneurysms, will be invaluable for establishing a causal relationship among hemodynamic stresses, inflammation and aneurysm growth.

Management of unruptured intracranial aneurysms still remains controversial. In some patients with unruptured aneurysms, the risk of surgical treatment may be higher than the risk of rupture. There are limited treatment options for some patients with giant aneurysms. New treatments targeting vascular remodeling and inflammation through inhibition of MMPs or other cytokines may be offered to these subsets of patients.

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REFERENCES
Intracranial aneurysms: links among inflammation, hemodynamics and vascular remodeling. T. Hashimoto et al.

54 Sadamasa N, Nozaki K, Hashimoto N. Disruption of gene for inducible nitric oxide synthase reduces progression of cerebral aneurysms. Stroke 2003; 34: 2980–2984

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84 Hashimoto T, Mesa-Tejada R, Quick CM, et al. Evidence of increased endothelial cell turnover in brain arteriovenous malformations. Neurosurgery 2001; **49**: 124–131
87 Achrol AS, Pawlikowska L, McCulloch CE, et al. Tumor necrosis factor-alpha-238G>A promoter polymorphism is associated with increased risk of new hemorrhage in the natural course of patients with brain arteriovenous malformations. Stroke 2006; **37**: 231–234