INTRODUCTION

Most unruptured aneurysms, which are detected incidentally show stable clinical courses, exhibit fewer inflammatory or degenerative changes in the walls of affected blood vessels, and have a low risk of rupture. However, some unruptured aneurysms show significant changes in size and shape, and the rupture risk seems to be high. These aneurysms may rupture in the early phase of development or enlarge in a short time due to thinning of vessel walls resulting from the advancement of degenerative changes. It is of clinical relevance to accurately estimate the rupture risk of cerebral aneurysms, but no definitive methods exist to distinguish rupture-prone aneurysms from rupture-resistant ones. Recently, the rupture risk of unruptured cerebral aneurysms was reported. The 5-year cumulative rupture rates for aneurysms located in the internal carotid artery, anterior communicating artery, anterior cerebral artery, or middle cerebral artery in patients without a history of subarachnoid hemorrhage were 0%, 2.6%, 14.5%, and 40% for aneurysms < 7 mm, 7-12 mm, 13-24 mm, and 25 mm or greater, respectively.[1] By comparison, the rupture rates of aneurysms involving the posterior circulation and the posterior communicating artery were 2.5%, 14.5%, 18.4%, and 50%, respectively, for the same size categories. In one study, the annual rupture rate of unruptured cerebral aneurysms in a Japanese cohort was 0.95%.[2] In another, the average annual risk of rupture associated with small unruptured aneurysms was 0.54% overall, 0.34% for single aneurysms and 0.95% for multiple aneurysms.[3] The molecular mechanisms leading to the occurrence, development, and rupture of cerebral aneurysms have been experimentally investigated. Ruptured aneurysms manifest significant endothelial damage, structural changes in vessel walls, and inflammatory cell invasion compared to unruptured aneurysms.[4] The walls of ruptured aneurysms are fragile, possibly because macrophage infiltration into the aneurysm wall results in the loss of smooth muscle cells and degeneration of matrix proteins. In this manuscript, we discuss the molecular mechanisms of cerebral aneurysm development, focusing on inflammatory processes.

INFLAMMATION AND ABDOMINAL AORTIC ANEURYSMS

The crucial role of inflammatory reactions can be seen in the formation of abdominal aortic aneurysms (AAAs). Important histological features of vessel walls with AAAs include chronic inflammatory cell infiltration of the adventitia and media, elastin fragmentation, degeneration, and attenuation of the media. Collagen in the media and adventitia provides tensile strength to the aortic wall. Collagen synthesis increases during the early stages of aneurysm formation, suggesting a repair process.[5] Inflammation-related mediators in aneurysm growth include matrix-degrading proteinases, proinflammatory cytokines, and chemokines.[6] In later
stages, degeneration of collagen exceeds its synthesis, and when accompanied by excessive degradation of other extracellular matrix macromolecules such as elastin, ultimately favors AAA rupture. Indeed, AAAs exhibit increased local production of enzymes capable of degrading the extracellular matrix proteins collagen and elastin.[7-9] Oxidative stress and elevation of hemodynamic stress lead to degeneration of elastin or collagen. The activation of Th1 cytokines via interferon gamma and interleukin-6 polymorphism is accelerated.[10,11] The combination of inflammatory reactions and inherited vascular fragility, along with environmental factors like advanced age or smoking, contributes to the formation and augmentation of AAAs.

MOLECULAR BIOLOGY OF CEREBRAL ANEURYSM FORMATION

Multiple inflammatory factors have been identified that play a crucial role in cerebral aneurysm formation.[12] Inflammatory cells such as macrophages, monocytes, and T lymphocytes have been found in aneurysm walls.[14,15] The infiltration of leukocytes is related to the impairment or elimination of collagen fibers. The plasma levels of cytokines, collagenase, and elastase are elevated in patients with cerebral aneurysms.[14,15] Recently, nuclear factor-kappa B (NF-κB) and tumor necrosis factor-alpha (TNF-α) have been widely investigated as potentially key molecules in the inflammatory process. NF-κB is a transcription factor that is known to be closely related to inflammation. NF-κB is activated in endothelial cells at the site of arterial bifurcation in the early stages of aneurysm formation, which induces hemodynamic stress.[16] This activation is attributable to hemodynamic stress in the affected endothelial cells. It is thought that activated NF-κB incites several downstream inflammation-related genes at the transcriptional level. Monocyte chemoattractant protein-1 (MCP-1) is one target of NF-κB and is an indispensable factor for the migration of macrophages to the lesion site. The transcription of MCP-1 is controlled by NF-κB at the cerebral aneurysmal wall. MCP-1 is secreted from the endothelial cell layer in the early stages of aneurysm formation and from all layers of the arterial wall in later stages. Macrophage infiltration can be suppressed by the use of MCP-1 knockout mice or MCP-1 inhibitor, and leads to inhibition of aneurysm formation.[17] This evidence highlights the significance of MCP-1 and macrophage infiltration. Other factors controlled by NF-κB are inducible nitric oxide synthase (iNOS) and interleukin-1 beta (IL-1β), which are known as apoptosis-inducing factors. An investigation of experimentally induced cerebral aneurysms in rats revealed that apoptosis occurs in smooth muscle cells located within the medial layer of affected vessels, and is associated with inflammation.[18] iNOS activity results in the production of nitric oxide, which is an important factor involved in inflammatory reactions and the preservation of arterial regulation. iNOS is principally expressed in inflammatory cells such as macrophages, and may impair arterial wall integrity or induce apoptosis. The expression of iNOS is facilitated in the media and adventitia during the early phase of aneurysm formation. The incidence of experimentally induced cerebral aneurysms in iNOS knockout mice is the same as that in control mice, however the aneurysm size is significantly smaller.[19] This suggests that iNOS contribute to aneurysmal augmentation by promoting apoptosis in medial smooth muscle cells. IL-1β is an inflammatory cytokine which is activated by cleaved caspase 1. IL-1β is also produced in the early phase of aneurysm formation, mainly by medial smooth muscle cells. In IL-1β knockout mice, the progression of aneurysm development is significantly impaired.[20] This means that inflammatory reactions in the arterial wall contribute to aneurysm enlargement and that IL-1β is a significant mediator of this process.

TUMOR NECROSIS FACTOR-ALPHA

Tumor necrosis factor-alpha has been revealed to have a close connection with several risk factors that affect aneurysm formation. Remarkable expression in aneurysm walls of mRNA for TNF-α has been observed in humans.[21,22] In addition, therapeutic administration of a TNF-α inhibitor significantly reduced aneurysm formation in rats.[23] There has been some investigation into the relationship between TNF-α expression and aneurysm formation or rupture. Inflammation induced by expression of TNF-α leads to the degeneration of endothelial cells, the internal elastic lamina, and medial smooth muscle. Cerebral aneurysms are stabilized when the expression of TNF-α is reduced, or expression of anti-inflammatory cytokines increases, however continuous expression of TNF-α induces aneurysmal rupture.[24] TNF-α also increases the permeability of the aneurysm wall via cytokine cascades and induces the migration of macrophages or neutrophils to inflamed endothelial cells. In addition, TNF-α plays a role in other pathological manifestations such as modulation of the blood-brain barrier, fluid accumulation, and regulation of intracranial blood flow in aneurysmal subarachnoid hemorrhage.[25] In transgenic mice that are deficient for TNF-α or TNF receptors, susceptibility to nitric oxide is notably increased.[26,27] In mice with congenital TNF-α receptor deficiency, the deleterious effects of oxidative stress are increased by traumatic or ischemic loading, which indicates that stimulation of antioxidant pathways by TNF-α may provide protection against oxidative damage.[28]
MECHANISMS OF CEREBRAL ANEURYSM FORMATION IN HUMANS

Aneurysm formation has been shown to occur at sites of constant hemodynamic stress both in humans and in experimentally induced models of cerebral aneurysm. This means that hemodynamic stress initiates early stage aneurysm formation. Aneurysm formation progresses when degenerative changes exceed vessel repair due to vascular remodeling. In addition to inflammation and apoptosis, extracellular matrix decomposition and endothelial dysfunction play critical roles in aneurysm formation. It has been found that gene expression in cerebral aneurysms may be linked to specific genetic regions. Furthermore, genetic regions that promote cerebral aneurysm formation are also linked to AAAs. Loss of vascular smooth muscle cells, thinning of collagen fibers, and inflammation are more prominent in the walls of ruptured aneurysms compared to unruptured ones. Therefore, these factors seem to play an important role in the weakening of aneurysmal walls [Figure 1].

CONCLUSION

Inflammatory cells have been found in the walls of cerebral aneurysms, and several inflammatory factors are reported to play crucial roles in cerebral aneurysm formation. Possible therapeutic interventions to reduce the formation of cerebral aneurysms may include the inhibition of mediators of inflammation.

REFERENCES


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