Hindsight

Linking endothelial dysfunction with endothelial cell activation

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The thin layer of cells that lines the interior of blood vessels, known as the endothelium, plays a complex role in vascular biology. The endothelium mediates blood vessel tone, hemostasis, neutrophil recruitment, hormone trafficking, and fluid filtration. Endothelial dysfunction, as defined by a lack of NO, has been linked to a variety of disease states, including atherosclerosis, diabetes mellitus, coronary artery disease, hypertension, and hypercholesterolemia. Indeed, restoration of endothelial function is one of the earliest recognizable benefits of statin therapy. In 1995, James Liao and colleagues published a study in the JCI demonstrating that NO is a vascular protective factor that limits endothelial activation and prevents leukocyte adhesion to the vessel wall.

Two converging concepts, endothelial dysfunction and endothelial activation, served as the basis for our 1995 JCI study (1). This study would not have been possible without the serendipitous contributions of our coinvestigators. At the time, we were interested in mechanisms underlying endothelial dysfunction, which is defined as the decreased synthesis, release, and/or activity of endothelium-derived NO, and how endothelial dysfunction could contribute to vascular disease. Gimbrone and colleagues were developing the concept of endothelial cell activation in response to nonlaminar flow as an early mediator of leukocyte trafficking in the vascular wall (2). Endothelial cell activation is defined by the endothelial expression of cell-surface adhesion molecules, such as VCAM-1, ICAM-1, and endothelial leukocyte adhesion molecule (ELAM, also known as E-selectin). Endothelial cell activation is typically induced by proinflammatory cytokines, such as TNF-α and IL-6, and facilitates the recruitment and attachment of circulating leukocytes to the vessel wall. In addition, Libby and colleagues were investigating signaling pathways that could lead to vascular inflammation and atherosclerosis (3). The close working relationship of our research laboratories provided us with the impetus to try to determine how endothelial dysfunction and endothelial cell activation were linked.

The idea that endothelium-derived NO could modulate leukocyte adhesion was first demonstrated by Kubes and colleagues, who showed that inhibitors of eNOS could increase neutrophil adherence to mesenteric venules (4). But the precise mechanism by which NO prevents leukocyte recruitment to the vascular wall was not known. To address this question, we first demonstrated that NO donors suppressed the expression of VCAM-1, ICAM-1, and E-selectin in response to a variety of proinflammatory cytokines. These results suggested a central mechanism for antiinflammatory effects of NO. Because the induction of these cellular adhesion molecules was mediated, in part, through the activation of the transcription factor, NF-κB (5), we postulated that areas of turbulent flow in the vessel wall, which would have less endothelium-derived NO, would be more prone to endothelial cell activation and atherosclerosis. Indeed, atherosclerotic lesions tend to develop in areas of vascular branching that are exposed to turbulent rather than laminar flow, and mice with targeted deletion of eNOS develop increased atherosclerosis and vascular inflammation (12).

Since the publication of our study, the known vascular protective role of endothelium-derived NO has expanded beyond its inhibitory effects on endothelial leukocyte interaction and leukocyte trafficking to include inhibitory effects on platelet reactivity, smooth muscle contraction and decreasing NO bioavailability through the induction of ROS (8). It was unclear how the endothelium was activated prior to the recruitment of inflammatory cells. A clue to this puzzle came from the seminal work of Li and colleagues, who showed that hypercholesterolemia brought on by an atherogenic diet could rapidly induce VCAM-1 expression in the rabbit endothelium (9). Additionally, hypercholesterolemia had previously been shown to produce endothelial dysfunction, in part, through inhibiting eNOS activity and/or NO bioavailability (10, 11).

Based upon these observations, we investigated whether endothelial dysfunction could lead to endothelial cell activation in the absence of cytokines. We found that suppression of basal eNOS activity by l-NG-arginine methylster (l-NAME) led to the rapid induction of VCAM-1 expression and increased monocyte adhesion (1), indicating that endogenous endothelium-derived NO could basally and tonically inhibit endothelial cell activation. These findings provided a mechanistic link by which mediators of endothelial dysfunction, such as hypercholesterolemia, smoking, or oxidative stress, could induce or enhance endothelial cell activation, leading to increased vasoconstriction, smooth muscle proliferation, platelet aggregation, leukocyte adhesion, LDL oxidation, and MMP activation (Figure 1). These results also suggested that areas of turbulent flow in the vessel wall, which would have less endothelium-derived NO, would be more prone to endothelial cell activation and atherosclerosis. Indeed, atherosclerotic lesions tend to develop in areas of vascular branching that are exposed to turbulent rather than laminar flow, and mice with targeted deletion of eNOS develop increased atherosclerosis and vascular inflammation (12).

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proliferation, and modulation of vascular matrix remodeling and angiogenesis. Conditions, such as laminar flow, or pharmacological agents, such as estrogens, HMG-CoA reductase inhibitors (statins), and angiotensin-converting enzyme inhibitors, which enhance endothelium-derived NO, have generally been found to be vascular protective and potentially antiatherogenic. In contrast, antiangiogenic therapies, which impair endothelium-derived NO through inhibition of VEGF signaling, occasionally cause endothelial dysfunction and hypertension in patients with cancer (13).

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