The Endothelium: Paracrine Mediator of Aortic Dissection

Running title: Seta et al.; Endothelial CypA Rips the Aortic Media

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For over 30 years the endothelium has assumed greater and greater importance in our understanding of the development of vascular pathology. This includes the discoveries that the endothelium releases powerful vasodilator and antiplatelet mediators, prostacyclin and nitric oxide, as well as its role in governing permeability, inflammation, and monocyte/macrophage infiltration of the blood vessel. In this issue of Circulation, Fan and colleagues show that boosting endothelial-derived oxidants in the mouse aorta by overexpression of the NADPH oxidase isoform, Nox2, during prolonged angiotensin II-induced hypertension, results in a high incidence of infra-renal aortic dissection.

Aortic dissections are often associated with aneurysms, which may affect both thoracic (TAA) and abdominal (AAA) regions, but can occur also in the absence of aneurysm. There are currently no approved drug treatments against these deadly aortopathies. Limited treatment options include blood pressure control, as a means of decreasing the risk of rupture, and endovascular or open surgical repair, a procedure with high risk of morbidity and mortality. Although the elucidation of the mechanisms responsible for early aortic pathological changes that precede overt dissection remains very challenging in humans, animal models suggest that medial degeneration, consisting of vascular smooth muscle cell apoptosis and elastin fragmentation, are early features. The work by Fan et al. identifies release from the endothelium of the pro-inflammatory cytokine cyclophillin A (cypA), upon angiotensin II administration, as a potential paracrine culprit mediating early degenerative events in vascular smooth muscle resulting in aortic dissection. Surprisingly, such a short-term dramatic effect was not a consequence of the hypertensive response to angiotensin II, as equal pressor doses of norepinephrine did not cause dissection. Instead, the authors present evidence that when Nox2 is over-expressed in the endothelium, angiotensin II causes excessive aortic inflammation and
remodeling leading to dissection because the augmented production of oxidants promotes the
secretion of cypA. The novel model they present, therefore, offers unique evidence of the
powerful influence of the endothelium on arterial wall structure as well as insights into potential
paracrine mediators within the arterial wall that may lead to aortic dissection. Despite the high
rate of dissection and the dramatic MRI pictures of the extent of the dissection along the aorta
presented by the authors, they did not report on any sudden deaths, the most dreaded
consequence of aortic dissection in man. Hopefully, further understanding of the molecular
mechanisms of dissection will provide therapeutic insights to prevent this cause of sudden death
in man.

Aortic dilatations and dissections, primarily in the thoracic region, are often a co-
morbidity of monogenic syndromes, such as Marfan’s, Loeys-Dietz’s and Ehlers Danlos’s
syndromes, characterized by genetic alterations in extracellular matrix components, including
fibrillin-1 and collagen. These gene mutations result in pathological aortic remodeling and
enlargement, which can progressively worsen into a thoracic aortic aneurysm. Similarly,
mutations in MYH11 and ACTA2, two major vascular smooth muscle structural proteins, result in
aneurysm. In addition, genetic mutations in transforming growth factor β (TGFβ)², a cytokine
with fibrotic effects on vascular smooth muscle, or TGFβ receptors (TGFβR1 and TGFβR2)³,
downstream effectors (SMADs)⁴ or inhibitors (SKI)⁵, have been implicated in the etiology of
aortic dilatation and dissection⁶. Losartan, an angiotensin II receptor blocker commonly used as
an anti-hypertensive medication, has recently entered clinical trials for the treatment of thoracic
aortic aneurysms in Marfan’s and related syndromes⁷, for its ability to antagonize the
downstream effector TGFβ⁸, as well as to decrease hemodynamic stress on the dilated aorta, by
lowering blood pressure. In contrast to purported deleterious actions in the thoracic aorta, TGFβ

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was protective in angiotensin II-induced AAA in normocholesterolemic mice by inhibiting inflammatory cell infiltration and consequent vascular smooth muscle cell apoptosis and extracellular matrix degradation\textsuperscript{9}. This underscores that molecular mechanisms may differ amongst the phenotypic spectrum of aortic aneurysms and dissections based on etiology and/or on location, making a “one-fits-all” therapeutic approach problematic. However, the fact that cypA mediates VSMC proliferation and recruitment of inflammatory cells in several arterial disease models including ligated carotid arteries, atherosclerosis and angiotensin II-induced aneurysms\textsuperscript{10} would suggest that cypA may be a common inflammatory mediator and a possible therapeutic target for several diseases of arterial remodeling characterized by oxidant overproduction and inflammatory cell infiltration.

Elastase infusions or calcium chloride instillation induce elastin fragmentation and mimic the structural defects of human aneurysmal aortas, and are also features of mice bearing mutation for genes required for structural integrity of the aortic wall, such as fibrillin-1, fibulin-4 and TGF\beta receptors. In addition, apolipoprotein E or low density lipoprotein receptor null mice administered angiotensin II have been extensively used as experimental models of aortic aneurysm, in part because of the reproducibility of the outcome, but despite the fact that hypercholesterolemia is not an independent risk factor for aneurysms in man. In the absence of the apolipoprotein E null genetic background, elevated doses of angiotensin II (commonly 3.2mg/kg/d) are required to induce aneurysms in mice and occur with a lower incidence than in hypercholesterolemic mice, although they cause similar tissue pathology. The relevance of angiotensin II-infusion models to the human pathology is still debated\textsuperscript{11}, largely because direct comparisons between animal and human specimens at early stages of the disease are not feasible. The striking finding by Fan et al. is that angiotensin II infusion, even at the modest dose of
1mg/kg/d in endothelial Nox-2 transgenic mice, was sufficient to cause dissections in 45% of their normolipidemic mice.

Angiotensin II infusion in rodents has been employed as a model of renin-dependent hypertension for more than 50 years. Despite the fact that the rapidity of development and severity of hypertension is not often reproduced in spontaneously hypertensive patients, the model continues to provide unique insights into the pathogenesis of hypertension and the mechanisms of its clinical sequelae. Two of the most exciting recent mechanistic observations provided by the model indicate the importance of NADPH oxidase in brain nuclei that control sympathetic nerve traffic, and the involvement of T-cells in mediating the hypertensive response to angiotensin II. These two new directions stem from the original fundamental observations that angiotensin II stimulates NADPH oxidase-derived oxidants and contributes to vascular constriction, hypertrophy and remodeling of the vascular wall, and atherosclerosis. The key NADPH oxidase involved contains the heme-binding subunit Nox2, or gp91phox, the isoform that accounts for superoxide anion production by neutrophils, macrophages, and other myeloid cells. A Nox2 deficient knockout mouse had diminished pressor and hypertrophic response to angiotensin II, but that study left open the question addressed in Fan et al. of whether Nox2 expression in different cell types was important. Interestingly, Nox2, even in the normal aortic wall, is concentrated in the endothelium as well as in adventitial fibroblasts, and these two sites also are where leukocytes increase during angiotensin II infusion. This inflammatory cell influx caused by angiotensin II is key as demonstrated by the fact that leukocyte infiltration as well as the pressor and hypertrophic response is diminished in a chemokine receptor knockout mouse. Fan et al. show that the increase in reactive oxygen species (ROS) that they have induced in the endothelium, leads to increased adhesion molecule
VCAM1 expression throughout the aortic wall, providing evidence that the greater inflammatory response due to a paracrine mediator is at the root of the increased incidence of aortic dissection.

Fan et al. also provide insights into the paracrine relationships within the vascular wall that mediate the response to angiotensin II. Earlier elegant studies showed that cypA in smooth muscle cells promotes inflammation and activation of proteolytic enzymes, and that mice doubly deficient in cypA and apolipoprotein E were prevented from developing aortic aneurysms during angiotensin II infusion. In a clever series of studies using conditioned medium of cultured endothelial cells and aorta from endothelial Nox2 transgenic mice, Fan et al. show that endothelial oxidants promote cypA production which in turn “primes” smooth muscle cells through Erk phosphorylation and increased oxidants. This, in turn, is responsible for the activation of proteolytic enzymes that destroy elastin and lead to dissection. The authors leave unaddressed the question of whether cell specific genetic deletion of Nox2 in aortic endothelium might prevent much of the aortic pathology in response to angiotensin II, which would further highlight the importance of paracrine mediators.

It is apparent that the paracrine relationships within the arterial wall induced by angiotensin II are multiple and complex, but that superoxide anion produced by NADPH oxidase, whether it be in endothelial cells, leukocytes, or adventitial fibroblasts is key. It is made clear by Fan et al. that endothelial oxidants can augment generation of cypA, but which oxidant species are involved and their cellular sites and enzymatic sources of origin is important to consider. In a supplemental figure, Fan et al. show that an inhibitor of nitric oxide synthase prevents much of the angiotensin II-induced ROS production by endothelial cells that overexpress Nox2, suggesting the possibility that uncoupled eNOS and the generation of the reaction product of superoxide anion and nitric oxide, peroxynitrite, is at work. Indeed, the
footprint of peroxynitrite, nitrotyrosine, is abundantly localized in the aortic intima and adventitia, but not in the media, after angiotensin II\textsuperscript{18}. In the media, evidence would suggest that hydrogen peroxide is the oxidant species at work, as angiotensin II-induced hypertrophy and aneurysm formation are prevented by smooth muscle specific overexpression of catalase\textsuperscript{21,22}. Either peroxynitrite or hydrogen peroxide can contribute, by transcriptional\textsuperscript{14} and post-transcriptional\textsuperscript{23} mechanisms, to the increase in matrix metalloproteinase expression and activation that was shown so well by Fan et al. to occur throughout the aortic wall and accounts for the elastin breaks and dissections they reported.

In considering multiple potential therapeutic targets to inhibit the molecular and cellular events leading to aortic dissection (Figure 1), it is important to realize that the same mechanisms that account for hypertension caused by angiotensin II, are clearly different from those which account for arterial hypertrophy and remodeling. For example, inhibiting the acute activation of NFκB by replacement of IκBα with IκBβ, nearly abolished the intense fibrotic response to angiotensin II in the aorta and heart without affecting the pressor response\textsuperscript{24}.

In addition, other agonists, such as endothelin, thromboxane A\textsubscript{2}, or cytokines such as TNFα can stimulate many of the pathways that angiotensin II does so potently and rapidly via the AT1 receptor, leaving the door open for potential trouble during the sole use of specific AT1 receptor antagonists. NFκB is clearly central to aortic pathologies, and yet its direct inhibition has been both elusive and considered unwise for the long term. Broad-spectrum agents to inhibit the inflammatory response, such as statins, are attractive though less specific possibilities\textsuperscript{25}. Making a target of the oxidants or their enzymatic sources, which are so clearly inherent to the mechanisms presented by Fan et al. has been a failure, indicated by the lack of conclusive evidence that antioxidants benefit cardiovascular disease. Directly targeting Nox2 chronically
has been deemed unwise because of its central role in leukocyte activation in infection. This potential side effect also has limited direct targeting of cypA which is implicated in a wide variety of pathologies in addition to aortic dissection, including cancer, viral infections, asthma, and rheumatoid arthritis\textsuperscript{10}. The most specific agent, cyclosporine, which binds directly to cypA, exerts profound immunosuppressive effects via its intracellular binding partner, calcineurin. There are agents in development that specifically bind to extracellular cypA avoiding these immunosuppressive effects\textsuperscript{10}. What would be useful are agents that effectively treat elevated blood pressure and, at the same time, prevent the long-term sequelae of vascular remodeling that accompany hypertension, so strikingly demonstrated in the study by Fan \textit{et al.}

CypA might also represent a novel biomarker of intramural degeneration preceding overt aortic aneurysm or dissection. Elevated serum or plasma levels of cypA are found in patients with unstable angina or after acute myocardial infarction\textsuperscript{26} and with stable coronary artery disease\textsuperscript{27}. Prospective studies on the temporal relationship between circulating cypA levels and the progression of aortic aneurysms or dissections are warranted to validate cypA as a diagnostic tool to identify patients more likely to benefit from long-term monitoring or immediate elective surgery.

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\textbf{References:}


**Figure Legend:**

**Figure 1.** Molecular mechanisms in the vascular wall leading to aortic dissection. In the presence of angiotensin II, endothelial Nox2-derived oxidants (O₂⁻ and ONOO⁻) stimulate endothelial cyclophilin A (cypA) production, which acts as a paracrine factor to activate metalloproteinases (MMPs) and ROS production in VSMC. MMPs, in turn, degrade elastin, causing aortic dissection. Angiotensin II elicits an array of oxidants (O₂⁻, ONOO⁻, H₂O₂) and inflammatory responses (NFκβ) within the arterial wall via AT1 receptors (AT1R), which stimulate VSMC and fibroblast proliferation and inflammatory influx in the vascular wall, all contributing to remodeling and fibrosis. Angiotensin II also stimulates cypA production in VSMC, further contributing to oxidants and inflammation.
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