Targeting von Willebrand Factor–Mediated Inflammation

Clive Drakeford, James S. O’Donnell

In this issue of *Arteriosclerosis, Thrombosis, and Vascular Biology*, Aymé et al1 provide new insights into the role played by von Willebrand factor (VWF) in modulating in vivo inflammatory responses. In particular, a single-domain antibody against the VWF A1 domain is shown to markedly attenuate leukocyte recruitment and vascular permeability in 2 distinct murine models of inflammation. Collectively, these findings support the hypothesis that VWF is involved in regulating inflammation and suggest that novel VWF-targeted therapies may be useful inhibitors of this critical step in inflammatory pathogenesis.

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VWF circulates in plasma as a large multimeric glycoprotein and plays critical roles in normal hemostasis.2 At sites of vascular damage, VWF binds to exposed subendothelial collagen. Shear stress then triggers unwinding of the normal globular conformation of VWF, leading to conformational changes within the A domains and in particular exposure of the GPIbα (glycoprotein Ibα)-binding site within the A1 domain. Consequently, tethered VWF can recruit platelets to the site of injury. In addition, VWF also acts as a carrier for procoagulant FVIII, protecting it against premature proteolytic degradation and clearance.3

Besides these important roles in maintaining hemostasis, recent studies have identified novel roles for VWF in modulating inflammatory responses.4–8 In vitro studies demonstrated that immobilized VWF binds directly to both polymorphonuclear leukocytes and monocytes under static and flow conditions.4 Under flow conditions, this VWF-mediated interaction involved transient rolling (mediated in part through VWF binding to leukocyte P-selectin glycoprotein ligand-1) followed by stable adhesion (mediated in part through VWF interaction with leukocyte β2-integrins). VWF binding to P-selectin glycoprotein ligand-1 was dependent on the A1 domain, whereas several regions of VWF (including D’D3 and A1A2A3) were implicated in β2-integrin binding.4

Roles for VWF in modulating inflammatory responses have also been observed in vivo.5–8 In an experimental sepsis model involving cecal ligation and puncture, Lerolle et al5 observed significantly enhanced overall survival in VWF−/− mice compared with wild-type controls. Importantly, this A1 domain single-domain llama-derived antibody cross-reacts with both human and murine VWF. Consequently, a bivalent variant of the single-domain llama-derived antibody (KB-VWF-006bi) interfered with ristocetin-induced murine thrombosis model. In keeping with previous studies, KB-VWF-006bi was shown to markedly reduce leukocyte extravasation in a murine model of immune complex–mediated vasculitis.7 Interestingly, in contrast to the critical need for platelet GPIbα in regulating VWF-induced neutrophil extravasation into inflamed peritoneum, Hillgruber et al7 showed that VWF-modulated neutrophil recruitment in cutaneous inflammation was GPIbα-independent.

To further elucidate the role of VWF in regulating inflammation in vivo, Aymé et al1 have developed a novel single-domain llama-derived antibody or nanobody that recognizes an epitope located within the A1 domain of VWF. Importantly, this A1 domain single-domain llama-derived antibody cross-reacts with both human and murine VWF. Consequently, this critical role for the A1 domain plays a specific role in facilitating the proinflammatory effects of VWF. Perhaps unsurprisingly, this critical role for the A1 domain is at least partially dependent on the presence of platelets. Cumulatively, these findings suggest that VWF binding to GPIbα serves to tether platelets, which, in turn, modulates endothelial cell barrier wall permeability and thus leukocyte tissue extravasation into the tissues.

These emerging data demonstrate that VWF influences multiple different aspects of inflammation in vivo (Figure). Further studies will be essential in defining the molecular mechanisms through which these VWF-mediated immunomodulatory effects are mediated. On the basis of the current
evidence, VWF clearly modulates inflammation through platelet-dependent and platelet-independent pathways. Moreover, although the underlying biology remains unexplained, the animal data further suggest that the relative importance of these discrete VWF-mediated pathways varies between different types of inflammation. Further insights into the role played by VWF in regulating leukocyte extravasation and endothelial cell permeability and the complex cross-talk that exists in vivo between hemostasis and inflammation will undoubtedly emerge in the near future. Given the significant morbidity and mortality associated with inflammatory pathology, defining the roles of VWF may offer exciting opportunities to develop novel therapies to address a critical unmet clinical need.

Disclosures
None.

References
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*Arterioscler Thromb Vasc Biol.* 2017;37:1590-1591
doi: 10.1161/ATVBAHA.117.309817

*Arteriosclerosis, Thrombosis, and Vascular Biology* is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
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Print ISSN: 1079-5642. Online ISSN: 1524-4636

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