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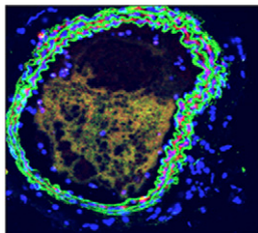
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molecules and NK cell receptors, suggesting that these two different types of interactions may perform distinct functions.

Provoking Platelets

Platelets express TLR4 and participate in sepsis following this receptor's stimulation by LPS. Because TLR4 and the receptor for the proinflammatory cytokine IL-1 act through common intracellular signaling components, Brown et al. (p. 5196) investigated mechanisms by which platelets might produce and/or be activated by IL-1. Human platelets were found to express IL1R1, but not IL1R2, and IL-1 β acted on IL1R1 to promote its own production via an autocrine loop. The active



IL-1 β produced by platelets was associated with microparticles shed by these cells. Similar to the action of LPS, IL-1 β stimulation led to activation of MyD88 and resulted in atypical platelet activation that included a failure to stimulate degranulation. Effective LPS-mediated stimulation in both human and mouse platelets was found to require signaling through IL-1R1 and splicing of pro-IL-1 β heteronuclear RNA to its active form. Using a model of arterial thrombosis to visualize activated platelets in vivo, the authors found that IL-1 β accumulated over time in platelets immobilized in occlusive thrombi. This accumulation did not require de novo RNA synthesis, suggesting IL-1 β production occurred by the mechanism of posttranscriptional RNA splicing observed in vitro. These data connect thrombosis to the development of chronic inflammation and reveal a mechanism by which IL-1 β and LPS work together to activate platelets.