Selectins revisited: the emerging role of platelets in inflammatory lung disease

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Commentary

Neutrophil infiltration into the lung is considered a crucial step in the pathogenesis of acute lung injury, yet data on the underlying mechanisms have been ambiguous: although selectin-mediated leukocyte rolling is absent in lung capillaries, therapeutic strategies targeted at selectin-mediated cell-cell interactions yield partial protection. The study by Zarbock and coworkers in this issue of the *JCI* solves this apparent contradiction by identifying selectin-mediated platelet-neutrophil interaction as a critical step in the mutual activation of leukocytes and endothelial cells (see the related article beginning on page 3211). The emerging role of platelets may be of broad clinical relevance in lung inflammatory disorders, including asthma, chronic obstructive pulmonary disease, and cystic fibrosis.
Neutrophil infiltration into the lung is considered a crucial step in the pathogenesis of acute lung injury, yet data on the underlying mechanisms have been ambiguous: although selectin-mediated leukocyte rolling is absent in lung capillaries, therapeutic strategies targeted at selectin-mediated cell-cell interactions yield partial protection. The study by Zarbock and coworkers in this issue of the JCI solves this apparent contradiction by identifying selectin-mediated platelet-neutrophil interaction as a critical step in the mutual activation of leukocytes and endothelial cells (see the related article beginning on page 3211). The emerging role of platelets may be of broad clinical relevance in lung inflammatory disorders, including asthma, chronic obstructive pulmonary disease, and cystic fibrosis.

Forty years after its first clinical description, acute lung injury (ALI) and its most severe form, the acute respiratory distress syndrome (ARDS), remain life-threatening conditions with reported age-adjusted incidences as high as 86.2 per 100,000 person-years and in-hospital mortality rates ranging between 31% and 38% (1, 2). The pathogenesis of ALI and ARDS is closely linked to intrapulmonary and systemic inflammatory responses. Neutrophils in particular have been implicated in the onset of both diseases based on the rapid accumulation of these cells observed in histologic lung specimens and bronchoalveolar lavage fluid from affected patients (3, 4). In many experimental models, a role for neutrophils in lung injury was supported by partial protection against injury following neutrophil depletion. Yet the causative role of neutrophils in ALI or ARDS has also been challenged by the clinical observation that even profound neutropenia does not protect patients from ALI and ARDS (5). Although the question of how many neutrophils are required to induce tissue injury remains unanswered, the latter observation suggests that neutrophil accumulation is not imperative for the onset of these diseases and that other inflammatory cells may be involved and compensate for neutropenia.

Neutrophil kinetics in the lung

In principle, neutrophil accumulation and subsequent tissue injury are the result of a multistep process comprising the initial tethering of circulating blood cells to the vessel wall and their subsequent rolling along the wall, followed by firm adherence and finally extravasation. This sequence of events is mediated by consecutive involvement of different families of adhesion molecules; while neutrophil rolling is mediated by selectins interacting with their respective glycoprotein counterligands, firm adherence results from the interaction of neutrophil β2-integrins with ICAMs expressed on the endothelium. In the systemic circulation, this sequence of events is predominantly confined to the venular compartment. In contrast, the prevalent site of leukocyte accumulation and emigration in the lung is the pulmonary capillary bed (6, 7). In lung capillaries, neutrophils do not roll but are temporarily retained at distinct sites of the alveolar capillary network for periods ranging from less than 1 second to more than 20 minutes (8). This phenomenon has been attributed to mechanical retention of circulating neutrophils in the narrow segments of the alveolar capillary network. Following mechanical arrest, the propelling blood flow slowly deforms neutrophils into an elongated shape, ultimately allowing them to continue their passage (9). In accordance with this notion, excessive accumulation of neutrophils in lung capillaries in systemic or pulmonary inflammation has been attributed to increased neutrophil stiffening by polymerization of monomeric to filamentous actin and subsequent firm adhesion to the endothelium via β2-integrins (10).
The role of selectins in neutrophil trafficking in the lungs was frequently considered negligible since the narrow pulmonary capillaries cannot accommodate the typical selectin-mediated rolling phenomenon. Furthermore, selectins did not seem necessary in lung neutrophil sequestration since the deceleration of circulating neutrophils prior to their firm adherence was effectively achieved by their mechanical retention. Yet the notion that neutrophils can accumulate in the lung and mediate ALI and ARDS without the need for selectin-mediated leukocyte rolling was strikingly inconsistent with a large body of experimental data demonstrating that selectin inhibition via the use of blocking antibodies or selectin antagonists or transgenic knockout of 1 or more selectins frequently protected animals from ALI (11). Thus, it seemed as if an important player linking neutrophil sequestration and selectin dependence in ALI still remained to be identified.

**Platelet-neutrophil interactions in ALI**

In an elegant combination of in vivo and in vitro experiments published in this issue of the JCI, Zarbock and coworkers unravel a critical role for platelets in the recruitment of neutrophils to the lung in experimental models of acid aspiration– and sepsis-induced lung injury (12). By platelet depletion, the authors attenuated lung histological changes, reduced protein leakage, and improved alveolar gas exchange in ALI. Importantly, platelet depletion also diminished the accumulation of neutrophils in all 3 compartments, i.e., the intravascular, the interstitial, and the alveolar spaces of the lung, demonstrating that platelet depletion interfered with an initial step of neutrophil accumulation and establishing a potential mechanistic link between platelets and neutrophils in ALI. P-selectin–dependent platelet-neutrophil interaction was identified as the structural nature of this link. By use of bone marrow chimeric mice, Zarbock et al. identified platelet-derived rather than endothelial-derived P-selectin as the relevant adhesion molecule mediating neutrophil sequestration and lung injury. It remains unclear whether platelet P-selectin interacts only with neutrophils or, in addition, also promotes platelet adhesion on lung endothelial cells, which would allow for “secondary capture” of neutrophils to the vascular wall (Figure 1). Recent intravital microscopic data suggest that activated platelets may attach to lung capillaries via platelet-derived P-selectin (13). This event may even precede the interaction of platelets with neutrophils because intrapulmonary causes of ALI, such as pneumonia or acid aspiration, can be expected to activate the vascular endothelium rather than circulating blood cells (14). Even systemic inflammatory stimuli seem to act primarily on the lung microvascular endothelium, as elegantly shown in experiments by Andonegui and coworkers, demonstrating that endotoxin-induced neutrophil accumulation is dependent on endothelial but not leukocytic expression of the lipopolysaccharide receptor TLR4 (15).

Following the initiation of neutrophil-platelet interaction, the reciprocal activation of both cells via outside-in signaling mechanisms accounts for the subsequent firm adhesion of neutrophils. In their in vitro experiments, Zarbock and colleagues (12) identified the eicosanoid thromboxane A2 (TXA2) as an important proinflammatory signal released by activated platelet-neutrophil aggregates, which mediates firm neutrophil adhesion by inducing the expression of endothelial adhesion molecules such as ICAM-1 (Figure 1). This notion of a critical role for a platelet-derived lipid mediator in ALI is consistent with the observation that transfusion of older, stored platelet concentrates containing bioactive lipids can cause transfusion-related ALI (16). It seems likely that other lipid mediators released from activated platelets, including platelet-activating factor and sphingolipids, contribute to ALI in a redundant or additive fashion. Yet the promising effects of both — a TXA2 receptor antagonist and a cyclooxygenase inhibitor — in the study by Zarbock et al. — render TXA2 and the eicosanoid pathway key candidates for intervention studies in ALI and ARDS. The failure of a previous clinical trial on the use of the thromboxane synthase inhibitor ketonocazole in ARDS does not conflict with this strategy since ketonocazole also failed to reduce the concentration of the stable TXA2 metabolite TXB2 in these patients (17).

**Platelets in inflammatory lung disease**

Over recent years, we have come to recognize the relevance of platelets not only in hemostasis, but also in numerous inflammatory processes, including atherogenesis, ischemia-reperfusion injury, and sepsis. Neutrophil-platelet interactions promote mutual cell activation, and platelet-endothelial interactions facilitate the secondary capture of neutrophils and other leukocytes. Zarbock and colleagues (12) have implemented this concept convincingly into the pathophysiology of ALI, but it may extend even further to other inflammatory respiratory diseases. Considerable numbers of circulating platelet-leukocyte aggregates can be found in patients with allergic asthma or cystic fibrosis (18, 19), and a critical role for platelet P-selectin in the recruitment of eosinophils and lymphocytes was recently demonstrated in an experimental model of allergic lung disease (20). Whether platelets alone may
In healthy individuals the immune system does not react aggressively toward host cells, a phenomenon defined as self tolerance. If self tolerance is broken autoimmune disease can develop, during which autoreactive lymphocytes are directed to a variety of autoantigenic epitopes. However, researchers have yet to determine whether immune responses to multiple autoantigens develop independently of each other or are the result of the response “spreading” from one autoantigen to another. In a study of NOD mice in this issue of the JCI, Krishnamurthy et al. show that the autoreactive T cell response to the autoantigen proinsulin lies upstream of that to islet-specific glucose-6-phosphatase catalytic subunit–related protein, suggesting that the pathogenic autoimmune response to proinsulin subsequently spreads to other antigens (see the related article beginning on page 3258). These data support the current view that this pancreatic β cell hormone is the first autoantigen targeted by the immune response in autoimmune diabetes.

Pathologic autoimmunity is characterized by an aberrant, self-perpetuating, immune-mediated, inflammatory response. It is the uncontrolled chronicity of this response that eventually leads to irreversible destruction of the target tissue. Among the major mechanisms underlying this chronicity is the diversification of the pathogenic autoimmune response, also termed epitope spreading.

The concept of epitope spreading was initially described by Eli Sercarz in the early 1990s in autoantigen-induced EAE, which is a model for multiple sclerosis (1). This term was used to describe how a self-directed immune response induced by a single peptide (or epitope) could spread to include other peptides (or epitopes) not only on the same autoantigen (i.e., intramolecular spreading), but also on other self molecules clustered in close vicinity within the target cell (i.e., intermolecular spreading). Thereafter, several studies confirmed the crucial role of epitope spreading in EAE (2–4) and also in demyelinating diseases of the central nervous system that follow some viral infections (e.g., Theiler’s murine encephalomyelitis; ref. 5) and IDDM, also known as type 1 diabetes (6, 7).