Expression of leukocyte integrins and tissue factor in mononuclear phagocytes

B. Nakstad*, T. Haugen*, O.H. Skjønsberg**, T. Lyberg*


ABSTRACT: Coagulation is intimately involved in the pathology of inflammation. The leukocyte β2-integrins have several functions, including serving as receptors for coagulation factor X and fibrinogen. Tissue factor (TF) is a receptor for factor VII and a very potent trigger of coagulation. The intention of this study was to examine a possible coregulation of β2-integrins (CD11b/CD18 and CD11c/CD18) and the procoagulant TF in alveolar macrophages (AM) and blood monocytes, i.e. cells of the same differentiation lineage.

The expression of β2-integrins in human AM isolated by bronchoalveolar lavage and in blood monocytes was analysed by flow cytometry, whereas TF activity was analysed in a one-stage clotting assay.

In monocytes, TF activity, CD11b and CD11c expression were highly inducible by lipopolysaccharide (LPS), with a 13-, 19- and four-fold increase, respectively. In AM, TF and β2-integrins were all constitutively expressed, but the expression could not be further enhanced by LPS stimulation. CD11b and CD11c expression varied inversely with the cell size of AM, in contrast to TF activity which is known to be proportional to AM cell size.

In vitro expression of β2-integrins and tissue factor in lipopolysaccharide-stimulated blood monocytes seems to be intimately coregulated, whereas the expression of these receptors in alveolar macrophages seems to be unresponsive to lipopolysaccharide. These results indicate that blood monocytes and alveolar macrophages have different roles and use different mechanisms in cell-induced fibrin formation.


The development of adult respiratory distress syndrome (ARDS), infant respiratory distress syndrome (IRDS) and septicaemia is known to be associated with pulmonary intravascular, interstitial and intra-alveolar deposition of fibrin [1]. The formation of fibrin during injury and inflammatory reactions, followed by its clearance during the repair process, is the result of a dynamic interplay between coagulant and fibrinolytic factors which amongst other cells are synthesized by peripheral blood monocytes, lung alveolar macrophages (AM) and lung interstitial macrophages [2–4]. Tissue factor (TF), which is produced by mononuclear phagocytes, is a receptor for coagulation factor VII [5] and plays an important role in the initiation of intravascular and extravascular coagulation leading to fibrin generation [2, 3]. Owing to their expression of strong procoagulant (TF) activity, monocytes and AM represent the major source of TF in the body [6] and in vivo [7, 8]. Binding of fibrinogen to specific receptors on blood monocytes has also been postulated to play a specific role in the differentiation of monocytes into macrophages [9]. Both monocytes and AM express β2-integrins (CD11b/CD18 and CD11c/CD18) on the cell surface which, in addition to mediating intercellular adhesion [10], have been shown to bind fibrinogen and factor X [11–13]. Coordinated membrane expression of procoagulant activity (TF) and receptors for coagulation factors (CD11b/CD18 and CD11c/CD18) would facilitate fibrin deposition when plasma factors and cofactors are supplemented. It is not known, however, whether these molecules are presented in a concerted fashion on the cell surface or whether they are independently regulated. Blood monocytes and AM are both members of the same cell differentiation (monocyte) lineage, but they are normally exposed to a different local milieu. It was, therefore, of interest to compare their response to the biologically highly active stimulant lipopolysaccharide (LPS), which in vivo is a potential candidate for cell stimulation during systemic pathological conditions such as septicaemia, but also locally when micro-organisms are introduced via the respiratory tract. LPS is a bacterial product of the outer membrane of Gram-negative bacteria which exerts its effects both directly, but also indirectly via the release of proinflammatory cytokines such as tumour necrosis factor (TNF)-α, interleukin (IL)-1β and IL-6. The intention of this study was, therefore, to compare the baseline expression of TF and β2-integrins on blood monocytes and AM and to examine the changed expression induced by LPS challenge.

Materials and methods

Bronchoalveolar lavage

Bronchoalveolar lavage (BAL) was performed in healthy, nonsmoking volunteers with normal lung function.
(seven males and seven females, aged 26±3 yrs (mean±SEM)). A Pentax FB 15X (Asahi Optica Co., Tokyo, Japan) bronchoscope wedged in a segmental bronchus of the middle lobe was used and 50 mL (×3) of sterile 0.9% NaCl (37°C) was instilled. The lavage fluid was recovered by gentle suction and immediately cooled on crushed ice. The study was approved by the regional ethics committee.

Alveolar macrophages

AM were separated from BAL fluid by centrifugation at 800xg for 7 min. Cells were then washed twice (300xg for 5 min) in phosphate buffered saline (PBS) and finally resuspended in endotoxin-free (<50 pg·mL⁻¹) cell culture medium (RPMI 1640; Gibco-Biocult, Paisley, UK) supplemented with 20% inactivated (56°C, 30 min) foetal calf serum (Bio-Wittacker, Walkersville, ML, USA) and seeded in 24-well tissue culture clusters (Costar, Cambridge, MA, USA) for TF analysis. Viability of cells was examined using the trypan blue exclusion technique. Differential counting was performed on cytopsin preparations (Shandon cyto Spin 2, London, UK) and stained with a modified Giemsa technique (DiffQuick, Merz&Dade, Harleco, Philadelphia, PA, USA). For flow cytometric analysis cells were suspended and cultured in polystyrene tubes.

Blood monocytes

Monocytes were obtained from healthy, nonsmoking volunteers (three males and four females, aged 28±2 yrs (mean±SEM)) by Lymphoprep (Nycomed Pharma, Oslo, Norway) centrifugation and adherence to tissue culture clusters as described in a previous paper [14]. Using this technique monocytes adhere to the plastic surface, whereas lymphocytes are nonadherent and are consequently removed by washing.

Experimental procedures

Cell cultures of AM and monocytes were incubated at 37°C in humidified air with 5% CO₂ for different periods (up to 24 h) in the absence and presence of LPS (lipopolysaccharide B from Escherichia coli 026: B6, Difco Laboratories, Detroit, MI, USA) (100 ng·mL⁻¹). The choice of sampling times was based on preliminary studies for each parameter with seven to 10 different time points ranging 0–72 h illustrating the maximal response. LPS concentrations ranging 1 ng·mL⁻¹–100 µg·mL⁻¹ were preliminarily analysed. The only concentration used in this study (100 ng·mL⁻¹) was eventually chosen because it gave a close to maximal LPS response in monocytes in vitro as regards all of the studied parameters and this LPS concentration is likely to be obtained in plasma in severe endotoxaemia.

Tissue factor assay

AM (3–6 ×10⁶ cells·well⁻¹) and monocytes (1.0–1.5 ×10⁶ cells·well⁻¹) were handled as described above and, after incubation, the cells were scraped off with a rubber policeman, washed twice in saline, resuspended in 0.5 mL Veronal-buffered saline (pH 7.35) and frozen at -20°C. After thawing and manual homogenization, AM and monocyte homogenates were tested for TF activity [15] and total protein [16]. Reference curves for TF were prepared by dilution of a standard human brain TF preparation [17] arbitrarily chosen to contain 100 U·mL⁻¹. TF activity was expressed as U·mg cell protein⁻¹.

Flow cytometry

An aliquot of 50 µL of AM suspension (1–4 ×10⁶ cells) or whole blood was incubated for 30 min in the dark with either 20 µL CD11b-specific (Becton Dickinson, Rutherford, NJ, USA) or 10 µL CD11c-specific (Sigma, St Louis, MO, USA), monoclonal fluorescein-conjugated antibodies. After erythrocyte lysis (in the case of monocytes), cells were washed twice with 2 mL PBS and fixed in 0.5 mL 1% paraformaldehyde. Cell suspensions with 10 µL PBS served as controls. β₂-integrin expression was evaluated using a FACSscan (Becton Dickinson) flow cytometer with an argon laser operating at 488 nm. Mean fluorescence intensity (MFI) was calculated by measuring 5×10⁴ (AM) or 1×10⁶ (monocytes) events, using the PC-Lysys 1.0 software program (Becton Dickinson). Relative linear mean fluorescence intensity (RLMFI) was measured in arbitrary units and given as the ratio between specific and nonspecific (autofluorescence) linear fluorescence. Comparisons between gated areas of large, granulated macrophages and smaller, less granulated macrophages, were made as illustrated in figure 1a.

Statistics

Results are expressed as mean±SEM. The Wilcoxon’s rank sum test and signed rank test of paired data were used for estimating significance. A p-value <0.05 was considered statistically significant.

Results

The purity of the AM preparations used in these experiments was 92±2%, whereas the proportion of lymphocytes was 6±2% and neutrophils was 2±1%. The purity of the monocytes in the monolayered cultures was >90% in every experiment, whereas lymphocytes represented the rest of the cell population. The viability of cells was always >90% (92±1% for AM and 95±1% for monocytes).

CD11b expression in blood monocytes and alveolar macrophages

Blood monocytes had a higher (two-fold, p<0.05) basal CD11b expression than AM harvested by BAL (figs. 2a, 3a). Furthermore, monocytes had a significant potential to increase their CD11b expression (19-fold, p<0.001) after LPS stimulation; the peak CD11b expression was observed 2 h (fig. 3a) after in vitro exposure to LPS. In contrast, AM could not be stimulated to increase their CD11b expression significantly above basal levels. Flow cytometric scatter plot-based separation (forward scatter versus
side scatter) of AM into subfractions (fig. 1a) showed that CD11b was most extensively expressed in the gated area R1 including AM which were smaller and less granulated than AM in the gated area R2 of larger size and higher granularity (p<0.02, fig. 1b).

**CD11c expression in blood monocytes and alveolar macrophages**

No difference in basal CD11c expression was observed between AM and monocytes (figs. 2b and 3b). As noted for CD11b, monocytes had a significant potential to increase their CD11c expression, i.e. a four-fold increase above their basal expression. The maximal CD11c expression appeared after 6–8 h (fig. 3b) of in vitro LPS stimulation. AM, however, could not be stimulated to increase their expression of CD11c. In fact, during in vitro culture the expression of CD11c in AM decreased (fig. 2b). Flow cytometric scatter plot-based separation (forward scatter versus side scatter) of AM into subfractions (fig. 1a) showed that CD11c was most extensively expressed in the gated area R1 including AM which were smaller and less granulated than AM in the gated area R2 of larger size and higher granularity (p<0.05, fig. 1c).
Tissue factor activity of blood monocytes and alveolar macrophages

Basal TF activity was significantly higher (three-fold, \( p<0.01 \)) in AM than in monocytes (figs. 2c and 3c) and monocytes had a highly significant (\( p<0.001 \)) potential to increase the TF activity after LPS stimulation (fig. 3c). Maximally, a 13-fold increased expression of TF was noted and the maximal activity of TF was observed after 6–8 h of \textit{in vitro} LPS stimulation. AM could not be induced to increase their expression of TF activity and during \textit{in vitro} culture their activity decreased.

Discussion

In the present study we demonstrated an inducible coexpression of procoagulant activity (TF) and the \( \beta_2 \)-integrin receptors CD11b/CD18 and CD11c/CD18 in human blood monocytes, whereas in AM a constitutive, noninducible synthesis and expression of all three molecules were observed. Both monocytes and AM synthesize TF [2, 3, 14] and are also able to produce and bind coagulation factor VII [3, 18, 19]. The complex of TF and factor VII initiates the cellular assembly of the prothrombinase complex [20], by which prothrombin is converted to thrombin [18]. The leukocyte \( \beta_2 \)-integrins CD11b/CD18 and CD11c/CD18 mediate a variety of functions which are upregulated when the cell is activated [11, 12]. At least one of these functions is associated with new epitopes that bind fibrinogen and coagulation factor X [21]. The binding of factor X to CD11b has been shown to be followed by a proteolytic activation step converting factor X to its active form Xa [13].

In contrast to monocytes, AM were fairly unresponsive to stimulation by LPS \textit{in vitro}. The ability of monocytes to increase their procoagulant activity after LPS stimulation is an important factor in septicemia and could explain most of the coagulation aberrations and thrombotic complications associated with this disorder. The ability of monocytes to increase rapidly the expression of CD11b and CD11c indicates that these cells are endowed with a fast integrin-associated system for pericellular fibrin formation in addition to the slower TF-driven system. Mutual interactions and interplay between these systems are highly probable, but the relative importance of each of these pathways is not known in detail. As demonstrated in other reports [22, 23], when monocytes are activated, CD11b/CD18 and CD11c/CD18 may be translocated from intracellular storage pools to the cell membrane, where they are expressed within a short time to reach maximal expression, as shown in this paper, within 2 h after LPS stimulation. Increased TF expression, in contrast, is dependent on \textit{de novo} protein synthesis and therefore, requires more time.

In contrast to monocytes, AM apparently lack both the fast and the delayed phase of LPS-induced procoagulant activities. This could be a result of loss of properties, \textit{i.e.} response to stimulation, in the process of differentiation from monocytes to AM. Different theories about the origin of AM have emerged. One theory considers them to be end cells, derived from monocytes and incapable of division in the steady state [24]. According to a second theory a bone marrow-derived cell population in the interstitium of the lung produces pulmonary AM by local division and maturation [25]. Lastly, a theory on monocytes maturing in the lung capillaries and final mitosis in the alveoli has been proposed [26]. However, irrespective of origin, but owing to their distribution to the pulmonary air-fluid interface, AM may be exposed to several inhaled stimulants that may prime and activate the cells, inducing a permanent, unresponsive state as indicated for CD11b, CD11c and TF responses after LPS stimulation \textit{in vitro}. Nevertheless, AM show a constitutive, high expression of both TF and CD11b, which means that pericellular fibrin formation may take place when the necessary coagulation factors become available. The procoagulant potential of AM may be considered as a defence mechanism which becomes operable when the alveolocapillary membrane permeability is increased to a degree where coagulation factors can leak from blood plasma to the alveoli.

There seems to be an inverse distribution of \( \beta_2 \)-integrins and TF activity between subfractions of AM which does
not fit the hypothesis of a closely regulated expression of these receptors. TF activity in AM has been shown to be inversely proportional to cell buoyant density [14], whereas the expression of CD11b/CD18 and CD11c/CD18 seems to be proportional to AM density, as judged both by AM distribution in a flow cytometric forward versus side scatter plot (this study) and by direct examination of Percoll-separated subfractions (T. Haugen et al., unpublished data). The skewed distribution of CD11b/CD18 and CD11c/CD18 amongst subfractions of AM may also reflect that monocytes turn on the expression of these adhesion molecules in the process of their migration from the vasculature to become AM and that the same molecules are downregulated during the further differentiation process of AM. The importance of CD11b-induced coagulation thus probably decreases as AM mature, while the role of the TF-driven system is maintained or even enhanced, in line with earlier observations that TF represents a differentiation marker for mononuclear phagocytes [27, 28] and, thus, TF is maximally expressed in the most mature AM.

The binding of factor X and fibrinogen to monocytes and AM is probably, to a large extent, mediated by CD11b/CD18 [11, 12]. However, our group has previously reported the presence of another integrin on mononuclear phagocytes [29]. The role of this integrin in mononuclear phagocyte-associated fibrinogen binding and fibrin generation is not yet settled. Still another integrin is interesting when coregulation of integrin receptors and procoagulant activity is discussed. It has been shown that ligand engagement of a β3-integrin (αI or βI) may induce high levels of TF expression in mononuclear phagocytes [30].

Alveolar fibrin deposition seems to be a prominent feature in diffuse alveolar damage, owing to the presence of active procoagulants [31] and the increased alveolectal leak of clotting factors [32]. Together with a persistent abnormality of fibrin clearance in the alveolar compartment during evolving diffuse alveolar damage [33, 34], this may explain the formation of hyaline membranes on bronchoalveolar surfaces in respiratory distress syndrome [35]. Fibrin, especially when it is crosslinked to fibronectin, and fibrinopeptides, provides a chemotactic stimulus for fibroblast recruitment as well as a positional matrix for fibroblast adhesion and replication [36, 37]. Fibroblast ingrowth into such fibrin deposits may have important consequences for the development of pulmonary fibrosis. In conclusion, the potential of monocytes and alveolar macrophages to stimulate fibrin deposition in diseased lung tissue may act indirectly as a stimulant for the development of pulmonary hyaline membrane deposition and fibrosis development in respiratory distress syndrome. In blood, lipopolysaccharide stimulation induces an acute upregulation of coagulation promoting receptors in monocytes, whereas within the respiratory lumen alveolar macrophages represent a constitutive and lipopolysaccharide-independent coagulation potential which comes into function when the permeability of the alveolectal membrane is increased.

References
20. Tracy PB, Eide LL, Mann KG. Human prothrombinase complex assembly and function of isolated peripheral


