Apoptotic human cells inhibit migration of granulocytes via release of lactoferrin

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Apoptosis is a noninflammatory, programmed form of cell death. One mechanism underlying the non-phlogistic nature of the apoptosis program is the swift phagocytosis of the dying cells. How apoptotic cells attract mononuclear phagocytes and not granulocytes, the professional phagocytes that accumulate at sites of inflammation, has not been determined. Here, we show that apoptotic human cell lines of diverse lineage synthesize and secrete lactoferrin, a pleiotropic glycoprotein with known antiinflammatory properties. We further demonstrated that lactoferrin selectively inhibited migration of granulocytes but not mononuclear phagocytes, both in vitro and in vivo. Finally, we were able to attribute this antiinflammatory function of lactoferrin to its effects on granulocyte signaling pathways that regulate cell adhesion and motility. Together, our results identify lactoferrin as an antiinflammatory component of the apoptosis milieu and define what we believe to be a novel antiinflammatory property of lactoferrin: the ability to function as a negative regulator of granulocyte migration.

Introduction
Apoptosis is a programmed, physiological form of cell death that, in inflammatory terms, is quiet: apoptotic cells are rapidly phagocytosed by their neighbors or by mononuclear phagocytes that are attracted by chemotactic factors such as lysophosphatidylcholine (1) and fractalkine (2) released by apoptotic cells. The rapid engulfment of apoptotic cells mitigates against the potential tissue-injuring and proinflammatory features of dead cells as exemplified by necrosis (3–6). Indeed, failed or delayed clearance of apoptotic cells can have detrimental inflammatory consequences, including the development of autoimmune pathologies (7–9). The mechanisms underlying the non- or antiinflammatory nature of the apoptosis program are not understood in detail. Antiinflammatory mediators such as TGF-β1 and IL-10 are known to be produced at sites of apoptosis, both directly by apoptotic cells themselves and indirectly through interaction of apoptotic cells with phagocytes (10, 11). Apoptotic cells selectively attract mononuclear phagocytes (1, 12), but, curiously, the other class of professional phagocytes, granulocytes or polymorphonuclear phagocytes, do not migrate toward apoptotic cells in vitro (12) and do not normally engage in apoptotic cell engulfment. The absence of granulocytes from sites of homeostatic and developmental apoptosis suggests that apoptotic cells have the potential to selectively regulate the recruitment of mononuclear leukocytes.

In marked contrast to sites of apoptosis, sites of acute infection are characterized by the presence of granulocytes, most commonly neutrophils, as a means to protect the host by engulfing, killing, and digesting invading infectious agents. Neutrophils, as a first line of immune defense, are rapidly recruited to the site of infection in response to a variety of inflammatory stimuli, including chemokines, cytokines, leukotrienes, and bacterial components such as LPS and N-formylated peptides (13, 14). Comparative studies on the slime mold Dictyostelium discoideum have shown that in response to chemotaxtants, neutrophils orient themselves and migrate in an “ameboid motion” by anterior pseudopod extension accompanied by simultaneous posterior contraction and retrac tion. Such polarized morphology is characterized by the formation of a lamellipodium at the leading edge and a uropod at the trailing edge of the neutrophil (15–18). This process is tightly controlled not only to ensure the efficient migration of neutrophils to inflammatory sites, but also to prevent their aberrant infiltration and consequent tissue-damaging activities. Detrimental effects of neutrophils — caused, for example, by release of their proteolytic enzymes — contribute to many pathological inflammatory conditions, ranging from vasculitis and ischemia/reperfusion injury to glomerulonephritis, rheumatoid arthritis, and acute graft rejection (19). Therefore, at inflammatory sites, negative signals exist that prevent neutrophil recruitment, dampen neutrophil responsiveness, and counterbalance or terminate the inflammatory response. This antiinflammatory program is characterized by cessation of neutrophil infiltration, as arachidonic acid–derived prosta glandins and leukotrienes are switched to lipoxins, resolvins, and pro protectins (20, 21). Resolution of inflammation is ultimately achieved through neutrophil apoptosis, and apoptotic neutrophils are subsequently phagocytosed by macrophages, a process that leads to the release not only of antiinflammatory cytokines but also of such antiinflammatory and proresolving lipid mediators as lipoxin A4, resolvin E1, and protectin D1 (22–25).

Given (a) the production of negative signaling molecules at inflammatory sites to limit neutrophil recruitment and function and (b) failure of neutrophil migration to sites of apoptosis, we sought to determine whether apoptotic cells actively produce negative regulators of neutrophil chemotaxis. We postulated that the production by apoptotic cells of factors that inhibit neutrophil migration contribute to the non-phlogistic nature of the apoptosis program. Here, we present evidence that apoptotic cells actively inhibit neutrophil migration through the production of...
lactoferrin, an 80 kDa antiinflammatory glycoprotein that specifically inhibits chemotaxis of neutrophils but not mononuclear phagocytes. Lactoferrin was found to be synthesized de novo following triggering of apoptosis and released by apoptotic cells of diverse lineages. Lactoferrin impaired neutrophil activation and prevented mobilization of the cells by inhibiting migratory polarization of the cell body. These results demonstrate that the non-phlogistic constitution of apoptotic cells includes an anti-inflammatory molecule, lactoferrin, that we show, for the first time to our knowledge, has potent negative regulatory effects on neutrophil migration. These findings provide a rationale for the absence of neutrophils from apoptotic sites and have important implications for understanding the mechanisms involved in the resolution phase of inflammation.

Results

Apoptotic cells actively produce factor(s) that inhibit neutrophil chemotaxis. To address whether apoptotic cells influence migratory activity of neutrophils, we carried out a series of in vitro Boyden-type chemotaxis assays to investigate neutrophil migration toward Burkitt lymphoma (BL) cells. We initially employed BL as a model system. As at all sites of apoptosis, there is marked infiltration of macrophages that engulf the apoptotic cells, giving rise to the typical “starry sky” histological appearance of this tumor. As shown in Figure 1A (left), while macrophages were abundant in histological sections of BL, neutrophils were absent. We assessed the effects of BL cells on the migratory activity of neutrophils in vitro by adding neutrophils to the top compartment of a Transwell filter and inducing them to migrate toward the lower chamber containing BL cells in the presence of the powerful neutrophil chemotactic formyl-methionyl-leucyl-phenylalanine (fMLP) (Figure 1B). As shown in Figure 1C, neutrophil migration was significantly inhibited in BL cells in a concentration-dependent manner. We observed a similar effect irrespective of the chemotactic stimulus used (inhibition of neutrophil migration induced by C5a, IL-8, and leukotriene B4 [LTB4]; data not shown, but see below). We carried out subsequent chemotaxis assays using BL-conditioned medium obtained over a 7-hour time course and found that BL cells actively released an inhibitory factor(s) (Figure 1D). The release of the inhibitory factor appeared to be linked to the levels of apoptosis in the BL cell populations, since the inhibitory activity was significantly lower in further chemotaxis assays using BL-conditioned medium derived from cells overexpressing the apoptosis inhibitor Bcl-2, as compared with that of parental cells (Figure 1E).

Biochemical characterization of neutrophil migration–inhibitory factor(s). In an attempt to gain further insight into the biochemical nature of the factor(s) that BL cells secrete in order to exclude neutrophils from their environment, we initially estimated the molecular weight range of the inhibitory factor(s) by using filters with molecular weight cutoff points of approximately 50, 10, 30, and 100 kDa. BL-conditioned media obtained after 24-hour incubation were fractionated, and each fraction was examined in vitro using the neutrophil chemotaxis assay described above. The results revealed that fractions containing molecules of less than 50 kDa failed to display any inhibitory effect on neutrophil migration (Figure 2A). However, the use of 100 kDa filters revealed that both fractions (>100 kDa and <100 kDa) displayed an inhibitory effect on neutrophil migration, indicating that at least 1 factor has a molecular weight that ranges between 50 and 100 kDa. The presence of inhibitory activity in the filtrate of the 100 kDa...
cutoff membrane is likely to result from (a) imprecise molecular weight cutoff of molecules in the 50–100 kDa range, (b) complex formation through multimerization of the 50–100 kDa factor or through interaction with other molecules, or (c) the existence of a distinct inhibitory activity of greater than 100 kDa. It should be noted that the selected isolation approach is skewed in favor of proteins and that additional low-molecular-weight (for example, lipid) mediators of neutrophil migration inhibition would not be identified by these procedures.

To investigate further the biochemical properties of the retentate and filtrate of the 100 kDa cutoff membrane, we first determined the charge (pI value) of the migration-inhibitory activity by means of an ion exchange analysis of BL-conditioned media. Using positively charged ion exchange beads (Q Sepharose), the 100 kDa membrane retentate and filtrate were separated into positively charged (Q-supernatant) and negatively charged (Q-eluant) fractions. As shown in Figure 2B, the filtrate of the 100 kDa membrane (<100 kDa fraction) contained migration-inhibitory activity in both the supernatant (positive charge) and the eluant (negative charge) of the Q beads. Neutrophil migration toward these fractions in the presence of fMLP (100 nM) was assessed. Q1 and Q2 fractions (unbound and eluant fraction) of serum-free medium (no BL) were included as control. *P < 0.05 compared with the corresponding control. Error bars indicate SEM. (C) Chemotaxis assay of neutrophils toward BL-conditioned medium that was heat inactivated (90°C for 10 minutes). (D) MALDI-TOF mass spectrum for the tryptic digest of the peptide band identified as lactoferrin.

Figure 2
Biochemical characterization of the inhibitory factor. Conditioned media from BL2 cells cultured for 24 hours were size fractionated using filters with 50 kDa (A) molecular weight cutoff sizes. Unfiltered medium was included as control. *P < 0.001 compared with the corresponding positive control. Error bars indicate SEM. Ion exchange analysis included the use of Q Sepharose beads (positively charged) in order to distinguish positively and negatively charged molecules in the <100 kDa fraction (B) of the BL medium. Unbound molecules (Q1 fraction) were collected, whereas bound molecules were eluted from the beads (Q2 fraction). Neutrophil migration toward these fractions in the presence of fMLP (100 nM) was assessed. Q1 and Q2 fractions (unbound and eluant fraction) of serum-free medium (no BL) were included as control. *P < 0.05 compared with the corresponding control. Error bars indicate SEM. (C) Chemotaxis assay of neutrophils toward BL-conditioned medium that was heat inactivated (90°C for 10 minutes). (D) MALDI-TOF mass spectrum for the tryptic digest of the peptide band identified as lactoferrin.
factor(s) were most likely protein in nature. We then analyzed the proteins released from BL cells in viable and apoptotic states by protein fingerprinting. Polypeptide bands of greater than 50 kDa were excised from a 10% SDS polyacrylamide gel. Tryptic peptides were gel extracted, and matrix-assisted laser desorption/ionization–time of flight (MALDI-TOF) mass spectrometric analysis was carried out (Figure 2D). Also, given the crude biochemical characteristics described above, we undertook a candidate approach based on the proteins released from apoptotic BL cells. We identified the factor released by BL cells that prevented neutrophil chemotaxis as lactoferrin.

Lactoferrin specifically inhibits neutrophil chemotaxis toward a range of chemoattractants. Lactoferrin is a glycoprotein of approximately 75–80 kDa that belongs to the transferrin family of proteins due to its iron-binding properties. It is a well-characterized component of neutrophil secondary granules, lacrimal fluid, colostrum, saliva, and mucosal secretions, in which it confers antibacterial activity. We observed that addition of anti-lactoferrin antibody to BL-conditioned medium neutralized its neutrophil migration–inhibitory activity using either polyclonal or monoclonal antibodies (Figure 3A and Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI36226DS1). Similar results were observed with supernatants obtained from the mammary carcinoma line MCF-7, indicating that the neutrophil migration–inhibitory activity is not restricted to BL cell–derived lactoferrin (Figure 3B). Furthermore, lactoferrin purified from human milk displayed dose-dependent inhibitory activity toward neutrophil migration in response to fMLP (Figure 3E) and also inhibited migration toward C5a, IL-8, and LTB4 to similar levels (Figure 4A). The neutrophil migration–inhibitory effect was also displayed by lactoferrin purified from human neutrophils (Figure 4D). It should be noted that both types of purified lactoferrin used in this study were free of endotoxin contamination, and the observed inhibitory effect did not appear to be due to any lactoferrin-associated molecules such as LPS. Furthermore, inhibition of lactoferrin expression by shRNA in BL cells provides additional support for the specificity of the observed lactoferrin effect. Thus, we found that neutrophil chemotaxis toward supernatants obtained from BL2 cells transfected with shRNA vectors targeted against lactoferrin was higher compared with supernatants from BL control or mock-transfected cells (Figures 3, C and D). Lactoferrin exerted no toxic effects on neutrophils, as assessed by annexin V/propidium iodide staining of control and lactoferrin-treated neutrophils (>98% cell viability). These results suggested that lactoferrin binds to neutrophils and inhibits their ability to undergo chemotaxis. To exclude the possibility that the observed inhibitory activity of lactoferrin was due instead to its ability to bind and functionally neutralize the chemoattractants, additional chemotaxis assays were performed using chemoattractants (fMLP, C5a, IL-8) that were preabsorbed with lactoferrin. To achieve this, we preincubated chemoattractants with lactoferrin. Subsequently, anti-lactoferrin antibody was used to remove the lactoferrin with the aid of magnetic beads. As shown in Figure 4B, no difference in neutrophil chemotactic activity was observed between the control and lactoferrin-absorbed chemoattractants, which excludes the possibility that lactoferrin binds to, and alters the activity of, the chemoattractants. Further supporting our conclusion that lactoferrin exerts its inhibitory effects through binding to neutrophils, we also observed that purified lactoferrin can directly associate with neutrophils (Figure 4C). In addition, Scatchard binding
analysis of $^{125}$I-labeled apolactoferrin indicated, in accordance with earlier findings (26, 27), that lactoferrin bound to neutrophils via 2 classes of receptor that differ in affinity and number of binding sites per cell. We determined the higher-affinity receptors to be expressed at a density of $9,100 \pm 2,500$ binding sites per cell, with an affinity of $350 \pm 65$ nM, and the lower-affinity receptors to be expressed at a density of $2.5 \times 10^6 \pm 0.7 \times 10^6$ per cell, with an affinity of $20 \pm 10$ μM (Supplemental Figure 2).

To determine whether the migration-inhibitory effects of lactoferrin were specific to neutrophils among professional phagocytes, we analyzed its effects on monocyte and macrophage migration in vitro. As shown in Figure 4, E and F, C5a-induced chemotaxis of mononuclear phagocytes was unimpaired by lactoferrin. We further assessed whether lactoferrin acted by inhibiting neutrophil migration or promoting neutrophil repulsion. In chemotaxis assays, in which we added lactoferrin to the upper chamber along with neutrophils, we observed inhibition of neutrophil migration toward fMLP and control medium (Figure 4G), suggesting that lactoferrin exerts a direct effect on neutrophils by inhibiting their migratory ability and not by forcing them to migrate in all directions away from the chemoattractant.

Neutrophil migration—inhibitory effects of lactoferrin are not related to its iron-binding properties and are demonstrable in vitro and in vivo. Iron and iron-associated molecules have been previously shown to play an important role in many immunomodulatory functions. Indeed, suppression of IL-1 release by monocytes is observed by purified iron-saturated lactoferrin, whereas an inhibition of GM colony-stimulating activity production by monocytes and macrophages correlated with the iron saturation status of lactoferrin (28–30). Therefore, we further examined whether differences in the iron saturation profile of lactoferrin affect its ability to inhibit neutrophil migration. Chemotaxis assays to determine neutrophil migration toward iron-depleted (apo-lactoferrin), partially iron-saturated, and fully iron-saturated (holo-) recombinant lactoferrin (10 μg/ml). Milk-purified lactoferrin and partially iron-saturated transferrin (TF; 10 μg/ml) were added as control. n = 4; *P < 0.001 compared with fMLP control. Error bars indicate SEM.
Lactoferrin inhibits neutrophil migration in vivo. Total cell (A) or neutrophil number (Gr-1+ B) obtained from peritoneal lavage. *P < 0.05 vs. transferrin; †P < 0.05 vs. thioglycollate (TG) control. **P < 0.01 vs. transferrin control. Error bars indicate SEM. (C) Characteristic cytospin images. Original magnification, ×400, top; ×200, bottom.

Impairment of neutrophil activation profile by lactoferrin revealed that the level of iron saturation was not responsible for the observed inhibition in neutrophil migration (Figure 4H). Also, as lactoferrin belongs to the transferrin family of proteins sharing 74% sequence homology with transferrin (both of them are 80 kDa cationic iron-binding glycoproteins), we reasoned that, if the underlying neutrophil migration–inhibitory mechanism of lactoferrin was rooted in its ability to chelate iron, transferrin might show similar effects on neutrophil migration. To explore this possibility, we performed chemotaxis assays in which neutrophils were induced to migrate toward fMLP in the presence of partially iron-saturated transferrin. Our results showed that transferrin, unlike lactoferrin, had no effect on neutrophil chemotaxis, further supporting the conclusion that the observed neutrophil migration–inhibitory effect is lactoferrin specific and is unlikely to require iron-chelating activity (Figure 4H).

Having established the inhibitory effects of lactoferrin on neutrophil chemotaxis in vitro, we then used a murine peritonitis model to assess the effect of lactoferrin on leukocyte recruitment in vivo. Lactoferrin and transferrin were tested for their ability to affect thioglycollate-induced leukocyte recruitment to the peritoneal cavity. As shown in Figure 5, A and B, thioglycollate caused rapid recruitment of leukocytes compared with vehicle alone, and the recruited leukocytes were predominantly neutrophils (88%). In the presence of lactoferrin, the total number of neutrophils recruited to the peritoneal cavity was reduced by 52% compared with control, whereas transferrin had no effect. Lactoferrin reduced specifically the proportion and number of neutrophils migrating into the cavity but did not affect recruitment of other types of leukocytes in response to thioglycollate (Figure 5C). These results demonstrate that, similar to its effect on neutrophil chemotaxis in vitro, lactoferrin is a potent inhibitor of neutrophil migration in vivo.

Neutrophil migration involves activation, adhesion, and extravasation processes that are accompanied by gross changes in cell morphology: whereas nonactivated neutrophils are rounded, activated neutrophils acquire a polarized morphology with spreading and adhesion to the available substratum (15). In order to initially assess the effects of lactoferrin on neutrophil activation, we performed time-lapse video microscopy of neutrophils and recorded directly their activation morphology, cell spreading, and locomotion. During a 1-hour time course, lactoferrin-pretreated neutrophil populations stimulated with fMLP displayed a greater proportion of nonadherent cells as well as cells presenting a rounded, nonactivated morphology as compared with neutrophils treated with fMLP alone (Figure 6, A and B). These quantitative differences between lactoferrin-treated and untreated neutrophils stimulated with fMLP were reflected in the locomotion of the cells around the substratum, with lactoferrin-treated cells displaying markedly reduced movement.

Changes in cell morphology following stimulation with fMLP or other neutrophil agonists are characterized by a rapid increase in intracellular cytoplasmic calcium levels through mobilization of calcium from ER stores and activation of calcium influx channels of the plasma membrane, mediated by the inositol triphosphate (IP3) and diacylglycerol/PLC (DAG/PLC) pathways (31–33). In order to determine whether the observed cell shape alterations following lactoferrin treatment are related to changes in intracellular calcium concentrations ([Ca2+]i), we measured the levels of [Ca2+]i in control and lactoferrin-treated cells in response to fMLP stimulation (1 nM or 10 nM). No changes were observed in the fMLP-activated [Ca2+]i response between lactoferrin- and control-treated neutrophils, suggesting that lactoferrin acts downstream or independently of the mechanisms involved in intracellular calcium flux (Figure 6C).

As lactoferrin was shown to prevent neutrophil migration, we next explored whether it could also affect the neutrophil activation state. To this end, we chose to measure the expression of 2
known neutrophil activation–associated markers, CD62L (L-selectin) and CD11b, using 2-color flow cytometry. Upon activation, CD62L is cleaved from the neutrophil surface, whereas CD11b expression is upregulated following translocation from cytoplasmic granules to the cell membrane. Freshly isolated neutrophils were pretreated with lactoferrin and then exposed to the activation stimuli fMLP, TNF-α, and PMA. As shown in Figure 7, A–D, we found that, in lactoferrin-treated neutrophils compared with control cells, CD62L expression was significantly higher, whereas CD11b levels were lower. These effects were common to all activation stimuli used. Transferrin-treated neutrophils were also included but showed no significant differences compared with control cells. It is noteworthy that the lactoferrin effect was also evident when PMA, a specific PKC activator, was used as an agonist, indicating that lactoferrin acts downstream of PKC and not on pathways involved in PKC activation and Ca\(^{2+}\) responses, such as the IP\(_3\) and DAG/PLC pathways. This finding prompted us to investigate putative downstream targets of PKC involved in the late signaling cascades following neutrophil activation that also regulate cell motility and actin reorganization. Such cascades involve the activation of MAP family kinases (34), and we therefore examined the phosphorylation of p44/42 (ERK1 and ERK2) MAPKs. Whereas in untreated neutrophils, ERK1 and ERK2 were phosphorylated following fMLP stimulation (100 nM), lower levels of phosphorylated ERK1/2 were observed in neutrophils that had been pretreated with lactoferrin prior to stimulation with fMLP (Figure 7E). Collectively, these data suggest that lactoferrin has a clear impact on neutrophil activation, including impairment of neutrophil degranulation, inhibition of expression of \(\beta_2\) integrins, and reduction of activation of intracellular kinases, with profound effects on cell migration and motility.

**Induction of apoptosis upregulates lactoferrin expression and release in diverse cell types.** Pursuing our initial hypothesis, which was strengthened by early observations that inhibition of neutrophil migration by BL cells appeared to be correlated with BL cell apoptosis (Figure 1E), we assessed lactoferrin expression following induction of apoptosis in a panel of cells of diverse lineages. By transcriptional analysis using RT-PCR, we found that lactoferrin was expressed, as reported previously (35), by MCF-7 mammary epithelial cells in their viable state but not by Jurkat, BL2, or A549 cells. Upon apoptosis induction, lactoferrin expression was upregulated in MCF-7 cells and expressed de novo in Jurkat, BL2, and A549 cells (Figure 8). More specifically, lactoferrin was transcribed de novo early after induction of apoptosis in A549 cells by either 100 nM etoposide or 1 \(\mu\)M staurosporine (Figure 8B). Levels of lactoferrin induced by etoposide were reduced in cells treated in the presence of the broad-spectrum caspase inhibitor zVAD-fmk, which prevented apoptosis induction (Figure 8C). The link between lactoferrin expression and apoptosis induction was further supported by the effects of the apoptosis inhibitor Bcl-2.
BL cells expressing exogenous Bcl-2 that provided protection from apoptosis expressed lower levels of lactoferrin upon exposure to staurosporine than did their parental counterparts (Figure 8, A and D). Not only was apoptosis-related lactoferrin expression demonstrated at the transcriptional level, lactoferrin protein was also recovered from supernatants of cells undergoing apoptosis (Figure 8D). Treatment of A549 cells with brefeldin A, which interferes with intracellular transport of newly synthesized proteins, resulted in inhibition of apoptosis-induced lactoferrin release, providing further evidence for de novo synthesis and secretion of lactoferrin by cells undergoing apoptosis (Figure 8E). An analogous effect was also evident from supernatants obtained from primary lymphocytes induced to become apoptotic in the presence of 1 μM staurosporine. Finally, immunoblotting analyses and chemotaxis assays using supernatants of BL cells undergoing primary necrosis further revealed that the release of lactoferrin is not linked to necrosis but that lactoferrin is expressed and actively released from cells as a consequence of activation of their apoptosis program (Figure 8F and Supplemental Figure 3).

Discussion

Apoptosis plays a key role in regulating cell populations through programmed cell death, an intrinsically non-phlogistic process that contrasts with accidental, necrotic cell death, which has proinflammatory consequences. Defining the properties of apoptotic cells that contribute to the noninflammatory or antiinflammatory nature of the apoptosis program is critical to our understanding of this fundamental biological process. We have now identified lactoferrin as a cell-autonomous antiinflammatory mediator that is produced by cells of diverse lineages undergoing apoptosis. Furthermore, these studies demonstrate a hitherto unrecognized immunomodulatory function of lactoferrin: to inhibit directly the migration of neutrophil granulocytes. Lactoferrin is therefore a constitutive antiinflammatory component of apoptotic cells that, in addition to its known antiinflammatory properties, militates against the proinflammatory recruitment of granulocytes to sites of apoptosis.

First discovered in milk almost 70 years ago, lactoferrin is a highly pleiotropic iron-binding glycoprotein closely related to transferrin that is a well-known constituent of exocrine secretions and secondary granules of neutrophils. It has many antiinflammatory and immunoregulatory properties, and the majority of these are not related to its ability to chelate iron, although some of its bacteriostatic features are dependent on iron binding (36, 37). Its ability to inhibit neutrophil migration, as described here, is an additional antiinflammatory feature of the molecule that is independent of its iron-chelating activity. In addition to its well-established antimicrobial properties, lactoferrin can inhibit proinflammatory responses not only through its ability to bind to key components of bacteria and viruses, but also through its direct activity on innate immune cells and molecules (36, 37). Thus, lactoferrin can inhibit production of proinflammatory mediators such as TNF-α and IL-6 (38, 39) and can promote production of antiinflammatory mediators including IL-10, IL-4, and TGF-β1 (40, 41).

The identification of specialized high- or low-affinity lactoferrin receptors on human neutrophils provides an initial mechanistic insight into how lactoferrin exerts its inhibitory effect on the neutrophil migration machinery. This is in accordance with our
results demonstrating that lactoferrin binds directly to neutrophils and impinges on intracellular signaling events coupled to neutrophil activation, rather than interacting with chemoattractants and modulating their activity. Moreover, the latter possibility was further excluded by the fact that the observed inhibitory effect of lactoferrin was (a) specific to neutrophil, not monocyte, migration, (b) observed with a diversity of chemoattractants, and (c) not directly neutralized by chemoattractants. Consistent with other studies (26, 27), our results demonstrated 2 classes of lactoferrin receptor on the surface of human neutrophils. Our results indicate that a relatively high-affinity receptor is expressed at relatively low density and that a relatively low-affinity receptor is expressed at relatively high density. However, the detailed characterization of the lactoferrin receptor(s) and dissection of downstream signaling pathways remain a significant challenge, as lactoferrin has been extensively reported to interact with a range of molecules and receptors in a cell-specific manner and therefore has the potential to elicit different signaling cascades in different cell types. Indeed, many researchers over many years have attempted the detailed characterization of lactoferrin receptors from various cell types with only partial success, and many candidate receptors have been proposed for different cell types. Importantly, as has been suggested elsewhere (42), critical differences in the identification of specific lactoferrin receptors stem from the cationic nature of the lactoferrin protein, allowing it to bind to anionic molecules found in virtually all cell types.

Here, we have observed several effects of lactoferrin on cell activation with clear implications for migration, adhesion, and motility of neutrophils. The initial phase of neutrophil activation following agonist stimulation is characterized by a rapid increase in \([\text{Ca}^{2+}]_i\), which in turn triggers the activation of PKC as well as a range of calcium-dependent signaling pathways that control key neutrophil effector functions, including degranulation, cytoskeletal rearrangements, and cell migration (32, 43–45). Our data clearly suggest that the observed lactoferrin-mediated inhibitory effects on neutrophil chemotaxis could be attributed to modulation of molecules other than those involved in the \([\text{Ca}^{2+}]_i\) response following agonist stimulation. In addition, analysis of the expression of early activation markers such as CD62L and CD11b following stimulation with diverse neutrophil agonists, including PMA, a specific synthetic intracellular activator of PKC, further suggests that lactoferrin acts downstream or independently of the PKC pathway, affecting signaling cascade components of the later phases of neutrophil activation. This notion was strengthened by our observation that lactoferrin inhibited the phosphorylation of p44/42 (ERK1 and ERK2) MAPKs, indicating that its action is mediated, at least in part, via the MAPK pathway, a key pathway downstream of PKC with a crucial role in the regulation of cytoskeletal rearrangement and cell adhesion (46–48). The previously reported specific interaction of lactoferrin with the calcium-calmodulin complex (\([\text{Ca}^{2+}]_i/\text{CaM}\) provides further insight into the potential intracellular signaling targets of lactoferrin (49). The \([\text{Ca}^{2+}]_i/\text{CaM}\) complex is formed following the increase in \([\text{Ca}^{2+}]_i\), and PKC activation and regulates the activity of a number of pathways involved in cell adhesion and migration, including the ERK1/2 kinases (50). Indeed, the use of specific inhibitors of calmodulin activity has been demonstrated to mimic the effect of lactoferrin — inhibition of neutrophil migration — reported here (51–53). Lactoferrin has been shown previously to bind to receptors on mononuclear phagocytes and to inhibit proinflammatory responses via NF-κB (39, 54, 55). Taken together with the evidence that apoptotic cells can suppress proinflammatory responses of mononuclear phagocytes and can elicit antiinflammatory responses by these cells (10, 11), our results indicate that it is now reasonable to suspect that these effects are mediated, at least in part, by lactoferrin produced by apoptotic cells. However, the involvement of the NF-κB pathway in the inhibition of neutrophil migration seems unlikely, since lactoferrin-mediated inhibition of chemotaxis of
neutrophils treated with gliotoxin, a specific NF-κB inhibitor, was found to be identical to that of untreated cells (data not shown). In addition, no changes in the levels of IkB, an inhibitor of NF-κB that is degraded following NF-κB activation, were observed in Western blot analysis of fMLP-stimulated neutrophils treated with or without lactoferrin (data not shown).

In general terms, in professional phagocytes, sensing mechanisms that are not yet well defined allow these cells to navigate to apoptotic cells through chemotactic processes involving lipid signaling molecules and classical chemokine mechanisms (1, 2). It is noteworthy that of the 2 categories of professional phagocytes — mononuclear (monocytes and macrophages) and polymorphonuclear (granulocytes) — apoptotic cells attract mononuclear phagocytes selectively (12). Indeed, the lack of recruitment of granulocytes to apoptotic zones is a hallmark of the non-phlogistic apoptosis program, distinguishing normal areas of cell death from sites of accidental or pathological tissue damage or infection. Recent evidence suggests that certain recombinant preparations of lactoferrin have the potential to chemotact monocytes (56). Although we failed to observe monocyte attraction to lactoferrin in our studies with milk-derived lactoferrin, these results indicate that lactoferrin has the potential to function in the selective attraction of monocytes but exclusion of granulocytes from sites of apoptosis. Our observation that lactoferrin, released into the supernatants of apoptotic cells, is able to inhibit the migration of neutrophils, demonstrates not only that physiological concentrations of lactoferrin are functional in mediating this effect but also that apoptotic cells are active in producing a factor that can potently suppress migration of neutrophils, the major subset of professional phagocytes among circulating leukocytes, to sites of apoptosis. Although it is clear that the secretory events inhibited by brefeldin A treatment (transport from the ER and Golgi network) are important for the release of lactoferrin from apoptotic cells, further work will be required to elucidate the details of this secretory process — including whether lactoferrin is transported directly from Golgi membranes for release from the cell surface.

While neutrophils may lack the sensing mechanisms to guide them to sites of apoptosis, the constitutive production of a factor that blocks neutrophil migration to such sites is likely to be important when the apoptosis program is activated in the presence of signals that promote neutrophil recruitment during inflammatory responses. In this context, it is noteworthy that lactoferrin is active in inhibiting IL-8 production by endothelial cells (57). Furthermore, lactoferrin’s ability to inhibit neutrophil migration is likely to be important in the resolution of acute inflammation. Thus, our results suggest that lactoferrin arising from the secondary granules of neutrophils and from apoptotic cells constitutes a negative feedback component that limits the influx of neutrophils to inflammatory sites. We would speculate that neutrophils undergoing apoptosis en route to resolution of inflammation would release lactoferrin as do other apoptotic cells but would be unlikely to be required to synthesize the protein de novo. The negative regulatory activity of lactoferrin described here places it as one of the few molecules, alongside lipoxins, netrin-1, and annexin-1, that negatively regulate neutrophil migration (21, 58, 59). More importantly, based on the high specificity of its migration-inhibitory properties to neutrophils, lactoferrin is identified here as a promising therapeutic target for a range of chronic inflammatory conditions, including vasculitis, pulmonary fibrosis, and ischemia/reperfusion injury, that are characterized by excessive neutrophil infiltration leading to neutrophil-mediated host tissue damage and remodeling (19).

Since the properties of lactoferrin extend well beyond its ability to regulate infection, immune responses, and inflammation, the link established here between lactoferrin production and apoptosis has many additional implications. First, lactoferrin’s ability to act as a growth factor (60, 61) extends evidence coupling the apoptosis program with tissue repair responses (62, 63). Second, additional recent evidence suggests that its protease activity may be instrumental in activating the caspase cascade. In particular, endogenous lactoferrin has been reported to induce apoptosis through activation of caspase-3 (64, 65). It is tempting to speculate that the de novo synthesis of lactoferrin by cells triggered to undergo apoptosis in our studies is linked to the initiation of the apoptosis program as well as to its antiinflammatory features. Therefore, lactoferrin may have multiple properties in apoptosis: (a) regulating the initiation of the program, (b) influencing repair in the tissue microenvironment, and (c) promoting the non-phlogistic nature of the process.

Lactoferrin’s functions in relation to tumor biology are currently unclear. Proteolyzed forms of exogenous lactoferrin have been reported to have proapoptotic and antitumor activities. In addition, lactoferrin appears to promote antitumor host immune responses (37), and activation of antigen-presenting cells has recently been reported using high doses of recombinant lactoferrin produced in Aspergillus (56). Its observed effects on cell growth are inconsistent, and both growth-promoting and growth-inhibiting effects have been reported (60, 61, 66, 67), suggesting the possible importance of tissue context. Since lactoferrin has potent ability to inhibit neutrophil migration, it seems likely that in tumors in which neutrophils play a supportive role, limitation of neutrophil infiltration through lactoferrin administration could be therapeutically beneficial. However, in tumors in which neutrophils are absent, we propose that, given the known antitumor effects of neutrophils (68–71), encouragement of neutrophil infiltration through inhibition of lactoferrin may effect tumor destruction. Our results might predict that tumors in which apoptosis is prominent would produce lactoferrin in situ. While this suggestion requires proper investigation, at least in one category, BL, a high-grade malignancy in which high-rate apoptosis occurs, in situ production of lactoferrin protein has been known for many years (72).

In conclusion, we describe a novel homeostatic function for the highly pleiotropic immunomodulatory glycoprotein lactoferrin. We demonstrate that lactoferrin production is closely coupled to the fundamental cell death program, apoptosis. In this respect, lactoferrin is much more generally expressed than previously realized. Our results show that lactoferrin endows apoptotic cells with antiinflammatory properties, including the capacity to inhibit neutrophil migration. Given the multifunctional abilities of lactoferrin, these results have broad implications for the influence of apoptotic cells on multiple physiological processes, including cell growth, differentiation, and innate and adaptive immune responses, as well as the pathological processes of inflammatory and malignant diseases.

Methods

Antibodies and reagents. The following antibodies and reagents were used in this study: rabbit polyclonal anti–human lactoferrin IgG antibody (Sigma–Aldrich), mouse monoclonal anti–human lactoferrin antibodies (LF-288, AbD Serotec; imab75 and imab77, ImmunoSolv), rabbit polyclonal immu-
noglobulin (IgG) negative control (Dako), IgG1 isotype control (Sigma-Aldrich), allopolytocyanin-conjugated (APC-conjugated) anti-CD11b (mlgG1; BD), FITC-conjugated anti-CD62L (mlgG1; AbD Serotec), purified human lactoferrin from milk (Sigma-Aldrich), purified human lactoferrin from neutrophils (Athens Research and Technology), recombinant human lactoferrin (Sigma-Aldrich), human recombinant apolactoferrin/ hololactoferrin (ProSpec), purified human transferrin (Sigma-Aldrich), recombinant human TNF-α (R&D Bioscience), PMA (Sigma-Aldrich), fMLP (Sigma-Aldrich), CsA (Sigma-Aldrich), IL-8 (Sigma-Aldrich), LTB4 (Sigma-Aldrich), brefeldin A (Sigma-Aldrich), staurosporine (Sigma-Aldrich), and giotoxin (Sigma-Aldrich). For lactoferrin absorption chemotactants, each chemotactractant was incubated with purified human milk lactoferrin (10 μg/mL), and a mouse monoclonal anti-human lactoferrin antibody (LF-2B8; 10 μg/mL) or isotype control was added. Antibodies were completely removed by BioMag goat anti-mouse IgG (QIAGEN) magnetic beads, and the efficiency of antibody removal was assessed by analysis of the preabsorbed chemotactants by anti-mouse IgG ELISA (data not shown). Furthermore, the depleted antibody was readily recovered from the beads (data not shown).

Cell isolation. Fresh human venous blood was collected from volunteers according to the Royal Infirmary of Edinburgh (Scotland) Research Ethics Committee (approval 1702/95/3/11), and mononuclear and polymorphonuclear (PMN) leukocytes were isolated as previously described (73). Neutrophils represented more than 95% of isolated PMN cells. Monocytes (>90% CD14+ cells) were positively selected from isolated mononuclear leukocytes using CD14 magnetic beads (Miltenyi Biotec). Human monocyte-derived macrophages were obtained following culture of monocytes for 6 days in Iscove’s modified Dulbecco’s medium (IMDM) containing 10% autologous serum.

Chemotaxis assay. In vitro leukocyte chemotaxis was measured following a well-established transfilter cell migration assay, as previously described (12) using polyvinyl uncoated Transwell inserts (5 μm pore size; Costar, Corning). Time of incubation (37°C; 5% CO2) varied for cell type (neutrophils: 60 minutes; monocytes: 90 minutes; macrophages: 4 hours). Unless otherwise stated, lactoferrin was used at 10 μg/mL. Chemotactic agents included fMLP (100 nM; Sigma Aldrich), CsA (6.25 ng/mL; Sigma Aldrich), IL-8 (50 nM; R&D Systems), and LTB4 (100 nM; Sigma Aldrich). For neutralization experiments, rabbit polyclonal anti–human lactoferrin antibody (Sigma-Aldrich) or negative control (Dako) were used. Filters were observed using an inverted microscope (Zeiss Axiosvert 25), and relative cell migration was determined by enumerating the number of migrated cells in 10 random high-power (×400) fields.

shRNA lentiviral transduction of BL cell. Lactoferrin expression was downregulated using shRNA lentiviral vectors (MISSION; Sigma-Aldrich). Briefly, lactoferrin-targeted shRNA lentiviral plasmids (pLKO.1-puro) were cotransfected with ViralPower Lentiviral packaging mix (pLIP1, pLIP2, pLP-VSV-G; Invitrogen) to 293FT cells using Lipofectamine LTX (Invitrogen). BL2 cells were transduced with the shRNA-expressing lentivirus, and stable cell lines were generated by selection with puromycin (2 μg/mL; Sigma-Aldrich).

RT-PCR analysis. Total RNA was extracted (RNeasy kit; QIAGEN) and reverse transcribed (2 μg) using SuperScript III RT (Invitrogen). Resulting cDNAs were used as template in PCR experiments at a concentration of 1 ng/50 μl of PCR mixture. The primers used were: forward lactoferrin (5’-TGCTTCTCTTGCTCTGTTCTC-3’) and reverse lactoferrin (5’-CTCCGCTCGATATGAACACCACATCA-3’), forward GAPDH primer (5’-CGACAGTACCCGACATTTCTTGGTCG-3’) and reverse GAPDH primer (5’-GGACTGGTTGCTGATAGCTTCACGAT-3’). Cycle parameters (lactoferrin: 40 cycles; GAPDH: 28 cycles) were: denaturation 94°C for 1 minute, primer annealing at 67°C (lactoferrin) or 50°C (GAPDH) for 1 minute, and extension at 72°C for 45 seconds, with a first denaturation step at 94°C for 7 minutes and a final extension at 72°C for 5 minutes. Purified PCR products (QIAquick gel extraction kit; QIAGEN) were sequenced to confirm validity by the Sequencing Service of the School of Life Sciences, University of Dundee, using Applied Biosystems BigDye 3.1 chemistry on an Applied Biosystems model 3730 automated capillary DNA sequence analyzer.

Peritonitis model. All animal procedures were carried out under a UK Home Office Animals (Scientific Procedures) Act 1986 project licence. Mice (8- to 12-week-old female C57BL/6 mice; n = 7 per group) were injected i.p. with purified human lactoferrin or transferrin (500 ng in saline/0.1% BSA Sigma Aldrich) or saline/0.1% BSA alone followed by a second i.p. injection with 1% thiglycollate (500 μl) or saline/0.1% BSA after 20 minutes. Recruited leukocytes were harvested after 4 hours by peritoneal lavage with ice-cold saline containing 2 mM EDTA. Harvested cells were counted using a NucleoCounter (ChemoMetec), which excluded nonnucleated cells. To determine the number of neutrophils (Gr-1+), cells were immunolabeled with PE-conjugated anti-mouse Ly6-Gr-1 and counted using Flow-Count beads (Beckman Coulter).

Histology and immunohistochemistry. Six- to 10-week-old BALB/c SCID mice were injected i.p. with 107 BL2 cells. Tumors developed i.p. within 2 months of injection. Mice were sacrificed and tumors excised. For positive control, BALB/c mice were immunized with sheep red blood cells and spleens harvested and frozen 7 days after i.p. injection. Immunohistochemistry was performed on frozen acetone-fixed sections (5 μm) of BL or spleen tissues using biotinylated anti–mouse Gr-1 antibody (10 μg/mL; BioLegend) or isotype control (AbD Serotec). Nonspecific adsorption of antibodies was blocked using serum-free Protein Block (Dako). Reactions were amplified using VECTASTAIN Elite ABC avidin-biotinylated peroxidase complexes (Vector Laboratories). Hematoxylin was used as counterstain.

Flow cytometry. Unless otherwise stated, cells were suspended in PBS containing 5% normal mouse serum or 0.1% BSA, and all antibody incubations were performed for 20 minutes on ice. Mouse neutrophils were defined based on the expression of Gr-1 epitope using PE-conjugated rat anti–mouse Ly6G (Gr-1; eBioscience). For the assessment of neutrophil activation, the following antibodies were used: FITC-conjugated anti-CD62L (F4C6, mlgG2b; AbD Serotec) and APC-conjugated anti-CD11b (ICRF44, mlgG1; BD). Isotype controls included mouse IgG1:FITC (AbD Serotec), mouse IgG1:APC (BD), and rat IgG2b:PE (eBioscience). Cell apoptosis was determined by labeling with annexin V and propidium iodide. Samples were analyzed using a BD FACSCalibur or FACScan cytometer, and data were analyzed using BD CellQuest software.

Size fractionation and ion exchange chromatography. Size fractionation of BL2 cell conditioned media was performed using filters with specific molecular weight cutoff sizes (Amicon Centrifugal filters YM-50 and YM-100; Millipore), following the manufacturer’s instructions. Ion exchange chromatography was carried out using Q Sepharose Fast Flow beads (Sigma-Aldrich). Beads were washed with neutralizing buffer, and bound proteins were then eluted by addition of 10 M NaAc, 0.5 M NaCl, pH 4. BL2 cell–conditioned medium or control medium (RPMI 1640) was mixed with the beads and incubated at room temperature for 5 minutes. Samples were centrifuged (300 g, 5 minutes) and supernatants stored. Beads were washed with neutralizing buffer, and bound proteins were then eluted by addition of the corresponding elution buffer (for S beads: 10 mM Tris, 0.5 M NaCl, pH 10; for Q beads: 10 mM NaAc, 0.5 M NaCl, pH 4). After a 5-minute incubation at room temperature, beads were centrifuged (300 g, 5 minutes) and supernatants collected and analyzed. Prior to chemotaxis analysis, the supernatants were diluted (1:100), and pH was adjusted to 7.0. Proteins were identified by peptide mass fingerprinting using MALDI-TOF mass spectrometry. The procedure was carried out by the Scottish Instrumentation and Resource Centre for Advanced Mass Spectrometry (SIRCAMS), School of Chemistry, University of Edinburgh.
Immunoblotting. Conditioned media from viable and apoptotic BL2 and AS49 cells were collected, and their protein content was TCA precipitated. Briefly, 100 μl TCA was added in 1 ml conditioned medium at 4°C. Samples were centrifuged at 18,000 g and the pellets washed in ice-cold acetone before resuspension in sample buffer (NuPAGE; Invitrogen). Neutrophils (5 × 10^6 cells/ml) were lysed for 15 minutes with 1% NP-40 in TBS containing protease inhibitor cocktail (Sigma Aldrich), aprotinin, leupeptin, pepstatin A, 4-(2-aminoethyl)benzenesulfonyl fluoride, sodium orthovanadate, benzamidine, levanosilone, and β-glycerophosphate. Samples were centrifuged (20,000 g, 4°C, 15 minutes) and resolved by SDS-PAGE using 4%–12% Bis-Tris gels (NuPAGE; Invitrogen). Proteins were then electrobotted onto a nitrocellulose membrane (NuPAGE; Invitrogen), blocked with 5% (v/v) for (lactoferrin) or 3% BSA (for ERK), and probed with monoclonal mouse anti-human lactoferrin (1:100; LF.28B; AbD Serotec) or mouse monoclonal anti-MAPK activated (diphosphorylated ERK1 and ERK2) antibody (1:1,000; Sigma-Aldrich) or polyclonal mouse ERK2 (1:1,000; Santa Cruz Biotechnology Inc.) or rabbit monoclonal anti-ikBα (74) (1:2,500; E130; Abcam) followed by HRP-conjugated goat anti–mouse IgG (1:2,000; Amersham) or HRP-conjugated goat anti–rabbit IgG (1:2,500; Dako) and visualized using ECL (GE Healthcare).

Binding studies. Biotinylation of human milk–derived neutrophils was performed using EZ-Link Sulfo-NHS-LC-Biotin (Pierce, Thermo Scientific). Fresh neutrophils were exposed to biotinylated lactoferrin (10 μg/ml) at 37°C for 1 hour, after which the cells were washed and lysates prepared, resolved by SDS-PAGE, electrobotted, blocked with 0.1% PBS-Tween, and probed with HRP-conjugated streptavidin (Dako). For Scatchard analyses, 125I-labeled human milk–derived lactoferrin, prepared using Pierce Iodination Reagent (Thermo Scientific), was added at a constant amount to fresh human neutrophils in the presence of increasing amounts of cold, unlabeled lactoferrin. After 30 minutes at 4°C, cells were washed 3 times before gamma counting.

Measurement of [Ca^{2+}]. Measurement of [Ca^{2+}], was performed as previously described (75). Briefly, freshly isolated neutrophils were resuspended (10^6/ml) in HBSS (without Ca^{2+}/Mg^{2+}) and were incubated with 2 μM FURA 2/AM (Calbiochem) at 37°C for 30 minutes. The cells were washed twice, resuspended at 2 × 10^6/ml in HBSS (with Ca^{2+}/Mg^{2+}), and incubated for an additional 30 minutes at 37°C in the presence of lactoferrin (10 μg/ml), and then stimulated with FMLP (1 nM or 10 nM). [Ca^{2+}], levels were determined based on 340-380 nm dual wavelength excitation in a PerkinElmer luminescence spectrometer at 37°C with constant stirring. Calibration was performed after each experiment using Triton X (Rmax) and EGTA (Rmax) and (Rmax) are the maximum and minimum 340-380 fluorescence ratios, respectively. [Ca^{2+}], was calculated based on the 340-380 nm fluorescence ratio.

Statistics. Results from multiple experiments are presented as mean ± SEM. One-way ANOVA was performed, followed by Bonferroni’s post-hoc test. In all cases, P values of 0.05 or less were considered significant.

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