In vivo expression of neutrophil inhibitory factor via gene transfer prevents lipopolysaccharide-induced lung neutrophil infiltration and injury by a beta2 integrin-dependent mechanism.

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In Vivo Expression of Neutrophil Inhibitory Factor via Gene Transfer Prevents Lipopolysaccharide-induced Lung Neutrophil Infiltration and Injury by a \( \beta_2 \) Integrin-dependent Mechanism

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Abstract

The binding of \( \beta_2 \) (CD18) integrins on PMN cell membrane to intercellular adhesion molecule (ICAM) counter-receptors on the surface of vascular endothelial cells mediates PMN adhesion to endothelial cells. Neutrophil inhibitory factor (NIF), a 41-kD glycoprotein isolated from the canine hookworm (Ancylostoma caninum), is a \( \beta_2 \) integrin antagonist that inhibits PMN adhesion to endothelial cells. We transferred the NIF gene into CD1 mouse lungs by intravenous injection of cationic liposomes to study the effects of in vivo NIF expression on LPS-induced lung PMN sequestration and the development of lung injury. RT-PCR and Northern blot analysis indicated the lung-selective expression of the NIF transgene, and immunocytochemistry showed prominent NIF expression in pulmonary microvascular endothelial cells. NIF staining was also observed in intraluminal leukocytes present in pulmonary microvessels. This may be the result of NIF binding to leukocytes after its secretion from the transduced lung cells, since there was no evidence of NIF gene expression in circulating leukocytes. Pulmonary vascular NIF expression abrogated the lung tissue PMN uptake and airspace migration of PMN and prevented lung vascular injury (as measured by the lung tissue uptake of \( ^{125}\)I-labeled albumin) after the intraperitoneal LPS challenge (200 \( \mu \)g/mouse). Expression of a control protein, chloramphenicol acetyltransferase (CAT), by the same strategy, had no effect on these responses. In vitro studies showed that NIF prevented mouse PMN adhesion consistent with the inhibition of lung uptake after LPS challenge in NIF transgene–expressing mice. We conclude that pulmonary vascular expression of NIF, a specific \( \beta_2 \) integrin–binding protein, is a potentially useful gene transfer strategy in modulating the infiltration of PMN across the alveolar-capillary epithelial barrier and in preventing lung vascular endothelial injury. (J. Clin. Invest. 1998. 101:2427–2437.) Key words: liposome-mediated gene transfer • CD11a • CD11b • CD18 • intercellular adhesion molecules

Introduction

Gram-negative sepsis is the most common cause of the adult respiratory distress syndrome, which is characterized by sequestration of PMNs in the pulmonary microcirculation and their migration into the alveolar space (1). LPS, a component of the wall of gram-negative bacteria, administered in vivo produces several pathophysiological changes characteristic of adult respiratory distress syndrome (1). Lung injury develops as the result of release of proteases and reactive oxygen species from PMNs adherent to pulmonary microvessel endothelial cells and infiltration of PMN into airspaces (2, 3).

The binding of CD11/CD18 or the \( \beta_2 \) family of integrins on PMN to the intercellular adhesion molecule (ICAM)\(^2\)-1 and ICAM-2 counter-receptors (3–5) expressed on the endothelial cell surface stabilizes PMN adhesion, and promotes PMN migration across the vessel wall after LPS exposure (3, 4). The \( \beta_2 \) integrins CD11a/CD18 and CD11b/CD18 are heterodimeric leukocyte surface receptors formed by the variant CD11 subunits and the common CD18 subunit (6–8). Expression of CD11a/CD18 and CD11b/CD18 integrins in response to proinflammatory cytokines and LPS contributes to leukocyte adhesion to the endothelium and both integrin family members mediate the adhesion-dependent degranulation and respiratory burst responses of PMN and coordinate their transendothelial migration (5, 9–11). Although CD11a/CD18 and CD11b/CD18 bind to several ligands, they bind specifically to ICAM-1 (CD54) and ICAM-2 (CD102) expressed on the endothelial surface (4, 12–15). It has been proposed that adhesion-dependent activation of PMNs signaled by the interaction of \( \beta_2 \) integrins with ICAMs is a critical event regulating the emigration of PMNs across the vascular endothelial barrier (3, 4, 13).

Studies have shown protective effects of both anti–CD11/CD18 and anti–ICAM-1 mAbs in models of acute lung injury (including that induced by septicemia) (5, 16, 17). Protection in these models has been universally attributed to inhibition of PMN sequestration in the lungs (5, 16, 17). However, the results using mAbs have not always agreed with experiments in mice made genetically deficient in adhesion molecules (5, 18–20). This may be the result of upregulation of compensatory pathways in response to the long-term loss of CD18 integrins or ICAM-1 in these knockout mouse models and to the potential nonspecific actions of mAbs, such as their ability to induce leukocytosis (21).

Neutrophil inhibitory factor (NIF), a glycosylated 41-kD protein isolated from canine hookworm (Ancylostoma caninum), is a potentially useful gene transfer strategy (including that induced by septicemia) (5, 16, 17). Protection in these models has been universally attributed to inhibition of PMN sequestration in the lungs (5, 16, 17). However, the results using mAbs have not always agreed with experiments in mice made genetically deficient in adhesion molecules (5, 18–20). This may be the result of upregulation of compensatory pathways in response to the long-term loss of CD18 integrins or ICAM-1 in these knockout mouse models and to the potential nonspecific actions of mAbs, such as their ability to induce leukocytosis (21).

1. Abbreviations used in this paper: BAL, bronchoalveolar lavage; CAT, chloramphenicol acetyltransferase; i.p., intraperitoneal(ly); i.t., intratracheal(ly); i.v., intravenous(ly); ICAM, intercellular adhesion molecule; MPO, myeloperoxidase; NIF, neutrophil inhibitory factor; rNIF, recombinant NIF protein; RT, reverse transcriptase.
the SuperScript Preamplification Reagent System (Life Technolo-
gen), inhibits PMN spreading and adhesion to endothelial
cells (22). NIF binds with high affinity to the metal ion-depen-
dent adhesion site on the A domain of the CD11b subunit of
PMN (22–24) and inhibits formyl methionyl leucyl phenylala-
nine (FMLP)–dependent adhesion of PMN to ICAM-1 and the
release of H2O2 by PMN (14, 22). We have shown that NIF
can also bind to CD11a expressed on activated PMN (Malik, A.B.,
and S.K. Lo, unpublished observation). In addition, we
showed that infusion of recombinant NIF protein (rNIF)
prevented lung PMN uptake and PMN-dependent lung injury
induced by TNF-α challenge of guinea pigs (25). Since the
in vivo expression of NIF may be a useful gene transfer strategy
in modulating or reversing lung PMN sequestration and the
trafficking of PMN into the airspace, in this study, we trans-
ferred the NIF gene into mouse pulmonary microvascular endo-
thelial cells using cationic liposomes and determined the ef-
effects of NIF expression on LPS-induced pulmonary PMN
sequestration and airspace migration of PMN, and on the
development of lung vascular injury.

Methods

Mice. Pathogen-free CD1 mice (6–12-wk-old males, 25–30 g body
wt) (Harlan Co., Indianapolis, IN) were used in all experiments. Mice
were housed in pathogen-free conditions in The University of Illinois
Animal Care Facility where they were treated in accordance with in-
stitutional and National Institutes of Health guidelines. Before injec-
tion of transgene/liposome constructs, the mice were anesthetized
with an intramuscular injection of ketamine (60 mg/kg) and xylazine
(5 mg/kg). The lipid molecules were sonicated for 20 min. The complex
described in formaldehyde and diluted (55°C for 15 min and elec-
trophoresed through formaldehyde-containing agarose gels. Northern
blot reagents were purchased from Stratagene (La Jolla, CA). Briefly,
RNA was blotted to Duralose-UV membranes (Stratagene) and the
membranes were cross-linked, hybridized, and washed as di-
rected by the manufacturer. The cDNA probes were labeled with
[a32P]dCTP (3,000 Ci/mmM) by the random priming method and the
membranes were exposed to Kodak autoradiography film (Eastman
Kodak Co., Rochester, NY) at −80°C for 6 h.

Histology and immunohistochemistry. Mouse lungs were inflated
with 4% paraformaldehyde in 1× PBS (pH 7.4) and embedded in
paraffin. Tissue blocks were sectioned to 5 μm thick and mounted
onto poly-L-lysine–treated glass slides. The lung tissue was stained
with hematoxylin and cosin to identify changes after NIF expression
in the intraperitoneal (i.p.) LPS-challenged mice (see below). For im-
munostaining, serial sections were incubated with blocking buffer
(1% goat serum in 1× PBS, pH 7.4) for 1 h at room temperature. A
rabbit anti–NIF polyclonal antibody (1:500) obtained from Dr.
Howard Soule was loaded onto sections and incubated overnight
at 4°C. Sections were then incubated with a biotinylated goat anti-rab-
bit IgG (1:250) in blocking buffer at room temperature for 1 h. Con-
trol sections were incubated with preimmune rabbit IgG and then, in
the same manner, with the secondary antibody (biotinylated goat
anti-rabbit IgG). All sections were incubated with colloidal gold-
labeled ExtrAvidin (1:50; Sigma Chemical Co.) for 10 min at room
temperature and counterstained with cosin after rinsing with PBS to
detect NIF expression.

LPS challenge. Except where otherwise indicated each mouse re-
ceived 200 μg of LPS (0111:B4; Sigma Chemical Co.) dissolved in 0.5
ml of PBS (pH 7.4) via i.p. injection. Control mice were injected i.p.
with an equal volume of PBS. Lung tissue myeloperoxidase (MPO)
activity was assayed (see Methods) to determine PMN sequestration
at different times after LPS injection (five mice in each group were
killed at each time point).

to determine the effects of NIF expression on LPS-induced lung
PMN sequestration, 5–10 mice in each group were injected i.v. with
PBS, liposomes, or transgene/liposome complexes 48 h before the
LPS challenge. Mice were then injected i.p. with LPS or PBS, and
were killed by a ketamine and xylazine overdose at 2 h, and the lungs
were removed for MPO activity, gene expression, bronchoalveolar la-
vage (BAL), vascular permeability, and morphological assessment.

BAL and leukocyte counts. The trachea was cannulated and
BAL was performed using 1 ml of PBS. BAL fluid was centrifuged
using a cytospin and BAL cells were stained with Diff Quik (Baxter
Healthcare, McGaw Park, IL). Differential cell counts were deter-
mined by counting 300 cells per slide (29). In some mice (n = 5 for
control and n = 5 for NIF transgene), LPS (25 μg) was instilled into
the trachea and BAL was carried out at 2 h as described above. The
circulating leukocyte count was determined by obtaining intracardiac
blood.

Lung PMN sequestration. Lungs were dried and homogenized in
0.5 ml of 50 mM PBS (pH 6.9) with 5% of hexadecyltrimethylamo-
nium bromide (ITAB) and 5 mM EDTA for quantification of PMN
uptake by MPO activity (30). The homogenates were sonicated, cen-
trifuged at 4 × 104 g for 20 min, and frozen and thawed two times fol-
lowed by homogenization and centrifugation. The supernatant was
mixed in 1:30 (vol/vol) with assay buffer (0.2 mg/ml of o-dianisidine
hydrochloride and 0.0005% H2O2), and absorbance change was mea-
sured at 460 nm for 5 min. MPO activity based on dry lung weight was
calculated as change in absorbance over time.

Lung PMN sequestration.
In Vivo NIF Expression Prevents Lung Injury

\[125I\text{-rNIF binding to mouse PMN.}\] rNIF (obtained from Dr. Howard Soule) was labeled with \(^{125}\text{I}\) (Amersham). \(^{125}\text{I}\)-rNIF was separated from free \(^{125}\text{I}\) by passage through a 10DG column (Bio-Rad Laboratories, Hercules, CA) in PBS containing 1% BSA as buffer. The specific activity of \(^{125}\text{I}\)-rNIF was \(\sim 8 \mu\text{Ci/\mu g}\). Mouse PMN (5 \(\times\) 10^5 cells) were incubated with 100 nM of \(^{125}\text{I}\)-rNIF on ice for 30 min with or without increasing concentrations of unlabeled rNIF. Mouse PMNs were isolated by the Ficoll gradient method and binding of \(^{125}\text{I}\)-rNIF was studied in PMNs challenged with LPS (10 ng/ml). The PMNs incubated with \(^{125}\text{I}\)-rNIF were washed with ice cold PBS and 1% BSA three times, and total cell-associated radioactivity was determined by \(\gamma\) counting (Auto-gamma 5000; Packard, Downers Grove, IL). Individual data points were obtained in triplicate to establish the binding of \(^{125}\text{I}\)-rNIF to PMNs. Competition studies were carried out to determine the effects of anti-CD11a and anti-CD11b mAbs added separately or in combination (PharMingen, San Diego, CA) on the binding of \(^{125}\text{I}\)-rNIF to PMNs.

In vitro mouse PMN adhesion. 8 h before harvesting PMNs, mice were injected i.p. with 2 ml of 4% Brewer’s thioglycolate broth (Difco Inc., Detroit, MI) (31). We chose this dose to obtain a maximal yield of PMNs. Prior experiments demonstrated that PMN yield was 80% of the total cells with the remaining amount being monocytes and lymphocytes. PMNs were shown to be viable by trypan blue exclusion. PMNs were harvested by peritoneal injection of 5–10 ml RPMI media to wash the peritoneal cavity and facilitate the removal of PMNs. Harvested cells were centrifuged at 1,000 rpm for 10 min. The total yield was 2 \(\times\) 10^7 cells per mouse. We determined the effects of NIF on the adhesion of mouse PMN to immobilized fibrinogen and to cultured human umbilical vein endothelial cells as described by Lo et al. (32).

Pulmonary vascular permeability index and wet/dry lung weight ratio. Mice were challenged with a 400-\(\mu\text{g}\) LPS i.p. injection 48 h after i.v. injection of liposome, pCR3/liposome complex, or pCR3-NIF/liposome complex. 3 h later, mice were i.v. injected with 1 \(\mu\text{Ci}\) of \(^{125}\text{I}\)-labeled albumin and killed 1 h after the injection of albumin tracer. Normal mice (i.e., those animals not challenged with LPS) were studied in the same way. Radioactivities of both blood and whole lung were counted to measure the pulmonary vascular permeability index (33) given to the ratio of lung counts per gram of dry lung weight-to-blood counts per gram of blood weight. The final wet/dry lung weight ratios in different groups were calculated to provide a measure of pulmonary edema.

Statistical analysis. Data are presented as mean \(\pm\) SEM unless otherwise indicated. Comparisons between experimental groups were made by ANOVA and the Wilcoxon test. \((P < 0.05).\)

Results

In vivo expression of NIF transgene and protein. The NIF gene and the reporter gene encoding CAT were separately constructed into the eukaryotic expression vector, pCR3, to transfer these exogenous genes in mouse lungs by liposomes. Liposomes were prepared and mixed with DNA constructs at a ratio of 1 \(\mu\text{g}\) of DNA to 8 nmol of liposome molecules (see Methods). The transgene (50 \(\mu\text{g}\)) was injected i.v. into each CD1 mouse in a total volume of 200 \(\mu\text{l}\) containing 5% glucose. The organs were removed at 48 h for gene expression analysis. Fig. 1, A and B, indicates the high and relatively selective CAT transgene expression in lungs after the i.v. injection of DNA/liposome complexes.

Lung NIF gene expression was initially determined by RT-PCR (Fig. 2 A). The DNA fragment covering the 720 bp...
NIF cDNA coding region was observed only in lungs of mice injected with the pCR3-NIF/liposome complex. This PCR product was not observed in control samples pretreated with DNase I and without reverse transcription, indicating the absence of DNA contamination in RNA samples. We observed, by immunostaining, the presence of NIF protein in pulmonary vascular endothelial cells and a dense staining pattern of the intraluminal leukocytes present in pulmonary microvessels (Fig. 2 B). In contrast, anti-NIF Ab did not stain red blood cells (which do not express β2 integrins) (Fig. 2 B). RT-PCR analysis of circulating leukocytes obtained from mice with a high level of NIF expression in their lungs showed that these cells did not express the NIF gene (Fig. 3). This observation is consistent with release of NIF into the pulmonary microcirculation resulting in its rapid binding to PMN (22, 23). In contrast to the leukocytosis evident after i.v. injection of anti-CD18 mAbs (5, 21), NIF expression by gene transfer in lungs did not raise the circulating leukocyte count (Table I).

125I-rNIF binds to mouse PMN and NIF prevents PMN adhesion. As the effects of NIF binding to LPS-activated mouse PMN are not known, we first determined the binding of 125I-labeled rNIF to isolated mouse PMN. The binding of 125I-rNIF to mouse PMN was specific and saturable (Fig. 4 A). Coincubation of 125I-rNIF with a 50-fold excess of unlabeled rNIF reduced the 125I-rNIF binding to PMN by 70%. Both anti-CD11a and anti-CD11b mAbs competed with 125I-rNIF binding to the LPS-activated PMN; each mAb decreased the binding of 125I-rNIF by ~40% (Fig. 4 A). Addition of both mAbs reduced the binding of 125I-NIF to PMN further to the same level (~70%) as observed with a 50-fold excess of unlabeled NIF (Fig. 4 A). NIF also prevented the adhesion of mouse PMN to immobilized fibrinogen and to endothelial cells in culture (Fig. 4 B).

In vivo expression of NIF in mouse lungs prevents LPS-induced PMN sequestration and lung vascular injury. All mice became lethargic and developed fur ruffling and diarrhea within 30 min of i.p. injection of 200 μg of LPS. 2 h after LPS injection, the MPO activity in lungs increased 3.5-fold compared to PBS-treated control mice (P < 0.05) (Fig. 5). PMN uptake remained high for 12 h, and returned to basal levels at 24 h after LPS challenge. Airspace PMN infiltration was a prominent feature of the lung pathology of the i.p. LPS-challenged mice (Fig. 6).

Mice were injected i.v. with the pCR3-NIF/liposome construct or control reagents (PBS, liposomes, wild-type pCR3/liposome, or pCR3-CAT/liposome complexes) to study the effects of pulmonary vascular expression of NIF in the i.p. LPS challenge model. Mice received an injection of either PBS or LPS i.p. 48 h after the injection of constructs (when significant NIF expression was evident), and were killed 2 h after PBS or LPS injection. The right lung was removed for gene expression analysis and the left lung for MPO activity. Analysis by Northern blotting showed NIF transcripts in lungs of both the PBS- and LPS-challenged pCR3-NIF injected mice (n = 10), but not in the control groups (n = 5 for each group) (Fig. 7). The lung MPO activity in LPS-challenged mice expressing NIF was 3.6-fold less than in control LPS-challenged mice (P < 0.01) (Fig. 8). The lung MPO activity in mice expressing NIF without i.p. LPS challenge was the same as in normal controls. CAT expression did not influence the increase in lung MPO activity induced by LPS (Fig. 8).

Mice became severely leukopenic within 2 h after i.p. LPS challenge (Table I), which is consistent with increased lung tissue uptake of PMN (Fig. 8). In contrast, the mice expressing NIF in pulmonary microvessels showed a modest decrease in the circulating leukocyte count (Table I), which is consistent with the observed inhibition of lung PMN uptake after i.p. LPS challenge of these animals (Fig. 8).

We also determined, by BAL, the effects of pulmonary vascular NIF expression on PMN migration into the alveolar space after i.p. LPS challenge. LPS challenge did not induce significant PMN infiltration into the alveolar space in mouse lungs expressing NIF, whereas control mice had markedly increased PMN numbers after similar intratracheal (i.t.) LPS challenge (Fig. 9). We compared these results with the effects
In Vivo NIF Expression Prevents Lung Injury

of pulmonary vascular NIF expression on airspace PMN migration induced by i.t. LPS challenge (Fig. 10). NIF expression reduced PMN migration in this model by ~ 50% (Fig. 10) (a significant inhibitory response \[P < 0.05\]) but less than the full inhibition of PMN migration observed in the NIF transgenic mice after i.p. LPS challenge [Fig. 9]).

The mice expressing NIF in their lungs, when challenged i.p., had exudate-free alveolar spaces, whereas lungs of control mice challenged i.p. with LPS showed prominent alveolar exudate in association with marked PMN infiltration (Fig. 6). Lung vascular permeability to albumin did not increase significantly in the NIF-expressing mice challenged with LPS,
whereas permeability increased two- to threefold in the control groups of mice (Fig. 11), indicating that NIF expression prevented the LPS-induced lung vascular injury. NIF expression also prevented the LPS-induced pulmonary edema as measured by the increase in the lung wet-to-dry weight ratio (Table II).

**Discussion**

LPS liberated from the outer membrane of gram-negative bacteria triggers PMNs to synthesize and release a cascade of inflammatory mediators that ultimately can lead to lung injury and multiple organ failure (1, 15, 34). LPS induces the expression of ICAM-1 and E-selectin on the endothelial cell surface and CD11/CD18 integrins on the PMN plasma membrane, which, through a series of binding interactions, mediates the adhesion of PMNs to the vascular endothelial monolayer and their migration across the endothelial barrier (5, 9, 10, 11, 35).

**Table II. Expression of NIF Prevents LPS-induced Increase in Wet/Dry Lung Weight Ratio**

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>LPS</th>
<th>LPS + liposome</th>
<th>LPS + pCR3</th>
<th>LPS + pCR3-NIF</th>
</tr>
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<tr>
<td><strong>n</strong></td>
<td>4</td>
<td>5</td>
<td>5</td>
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<td>5</td>
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<tr>
<td>Wet/dry ratio (Mean ± SEM)</td>
<td>4.7 ± 0.1</td>
<td>5.1 ± 0.1*</td>
<td>5.3 ± 0.1†</td>
<td>5.3 ± 0.1†</td>
<td>4.7 ± 0.1</td>
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*Values are mean ± SEM. †P < 0.01 compared with control group.

**Figure 3.** Leukocytes obtained from mice expressing the NIF transgene in lungs do not express NIF. NIF gene expression was determined by RT-PCR as described in Methods. From left: lane 1, DNA marker; lane 2, PCR from pCR3-NIF using NIF primers (positive control); lane 3, PCR using mouse leukocyte RNA (the mouse was injected i.v. with the pCR3-NIF liposome construct); lane 4, RT-PCR using mouse leukocyte RNA (the mouse was injected i.v. with the pCR3-NIF liposome construct); lane 5, direct PCR using lung RNA from the same mouse as lanes 3 and 4; lane 6, RT-PCR using mouse lung RNA from the same mouse as lanes 3 and 4. All RNA samples were pretreated with DNase I before PCR and RT-PCR. The results are representative of three separate experiments.

**Figure 4.** (A) Specific binding of 125I-rNIF to mouse PMN. Mouse PMNs (5 x 10⁵) activated with LPS (25 ng/ml) were incubated with 125I-rNIF in the absence or presence of unlabeled NIF, anti–mouse CD11a mAb, anti–mouse CD11b mAb, or both mAbs. Addition of increasing concentrations of unlabeled NIF reduced 125I-NIF binding by ~70%. Each mAb reduced 125I-NIF binding to mouse PMN ~40%, whereas the reduction with both mAbs was ~70%. Data are shown as mean ± SEM of three separate experiments for each concentration. (B) Mouse PMN adhesion to immobilized fibrinogen and human umbilical vein endothelial cells (HUVECs). Human fibrinogen was immobilized onto glass slides using HSA (1 mg/ml). NIF was added to the well before the adhesion assay was carried out for 10 min at 37°C as described previously (32). Mouse PMN adhered to the fibrinogen-coated surface after addition of phorbol 12-myristate 13-acetate (PMA; Sigma Chemical Co.), and NIF caused dose-dependent inhibition of PMN binding to fibrinogen. NIF also inhibited the PMA-activated adhesion of mouse PMNs to HUVECs. Data in both groups are shown as means ± SEM of three separate experiments in each group. *Difference from the untreated control group (–NIF) (P < 0.05).
In vitro studies showed that the stable adhesion of PMNs resulting from binding of CD11a/CD18 and CD11b/CD18 integrins to ICAM-1 and ICAM-2 expressed on the endothelial cell membrane regulates the transendothelial migration of PMNs (4, 35). Expression of CD11a/CD18 and CD11b/CD18 β2 integrins on PMNs mediates many of the proinflammatory functions of PMNs (5, 11, 35–37). In vitro studies showed that either anti-CD11a or anti-CD11b mAbs reduced PMN adhesion to endothelial cells by ~50%, whereas these mAbs in combination or mAbs directed against the common β subunit (CD18) fully prevented the PMN adhesion response (5, 32). Infusion of anti-CD18 mAbs also inhibited lung PMN sequestration in animal models (5, 16). However, the specific function of β2 integrin family members is unclear since mice made genetically deficient in CD11b did not show reduced PMN emigration across the endothelial barrier (11, 19).

In this study, we transferred the NIF gene, a recently cloned β2 integrin antagonist isolated from the canine hookworm (21, 22), to study the function of β2 integrins in regulating LPS-induced PMN trafficking in lungs and in the mechanism of lung vascular injury. We used cationic liposomes injected i.v. to express the NIF transgene and studied effects of in vivo NIF expression in an i.p. LPS challenge model that elicited lung tissue PMN sequestration and increased lung vascular permeability. We used i.v. injected cationic liposomes because this approach has been effective in inducing transient gene transfer in lungs (27, 38, 39). We showed that i.v. injection of CAT/liposome and NIF/liposome transgene constructs resulted in a lung-selective expression of these transgenes. NIF expression induced by this method was confined to pulmonary microvessel wall cells including the microvessel endothelial cells, consistent with recent evidence of endothelial cell uptake of cationic liposome–DNA complexes in mice (40). The selectivity of gene transfer in pulmonary microvessel cells may be related to factors such as the large pulmonary vascular surface area, delivery of the entire cardiac output to pulmonary circuit, and electrostatic charge-dependent binding of cationic liposomes to vascular endothelial cells (27). We also observed significant NIF immunostaining on membranes of leukocytes present in the lumen of pulmonary microvessels, perhaps reflecting the high affinity NIF binding to PMN β2 integrins (22, 23). The NIF gene was not expressed in circulating leukocytes of these mice, suggesting that positive NIF immunostaining of PMN is the result of the released NIF binding to leukocytes in transit through the pulmonary circuit.

In establishing the model of LPS-induced lung PMN infiltration, we showed that i.p. injection of LPS in mice resulted in a time-dependent increase in lung tissue MPO activity, which peaked 2 h after LPS challenge, and persisted for 12 h. Lung PMN sequestration in these mice was consistent with the severe leukopenia seen within 2 h after LPS challenge. We also showed an increased number of PMN recovered by BAL at 2 h after LPS challenge indicating the emigration of PMN into airspaces. Histological examination revealed PMN infiltration into both the interstitium and airspaces. These mice also showed alveolar exudate consistent with evidence of lung injury as described in CD1 mice challenged with LPS (41). The pulmonary vascular NIF expression in mice prevented LPS-induced leukopenia and lung PMN sequestration as well as the emigration of PMN into the airspaces. The in vivo NIF expression in lungs also prevented the LPS-induced increase in lung vascular permeability and edema formation.

The basis of the protective effects of NIF expression may be related to the binding of NIF to PMN β2 integrins, and the resultant inhibition of PMN adhesion. We showed that NIF was capable of fully inhibiting adhesion of mouse PMN. We also showed that LPS induced specific and saturable binding of 125I-labeled rNIF to mouse PMN and that anti-CD11a or anti-CD11b mAbs competed for 125I-NIF binding to these PMN. These results are consistent with our findings using phorbol ester–activated PMN and JY lymphoblastoid cells (which only express the CD11a/CD18 integrin) that NIF can bind to both CD11a and CD11b integrins on activated PMN (Malik, A.B., and S.K. Lo, unpublished observation). Therefore, a tenable explanation for the inhibition of PMN uptake in lungs expressing NIF may be that NIF binds to CD11a/CD18 and CD11b/CD18 integrins, thus, inactivating both integrins.

PMN migration into airspaces after the i.t. LPS instillation was inhibited only by ~50%, whereas pulmonary vascular NIF expression prevented the migration of PMN into airspaces after i.p. LPS challenge. This finding suggests that septecemia induces PMN migration across the microvessel and air-
way epithelial barriers via activation of β2 integrins. In contrast, airway gram-negative infection may induce PMN migration into airspaces by both β2 integrin-dependent and -independent mechanisms. Airway LPS may result in the release of chemokines (e.g., IL-8 [5] and macrophage inflammatory protein-1 [41]) capable of directing PMN into airspaces independent of CD18 integrins.

We evaluated the possibility that liposome injection may
In Vivo NIF Expression Prevents Lung Injury

have independently interfered with PMN uptake response and that this could account for our results. We showed that lipo-
some injection alone did not induce or prevent the increase in
lung PMN uptake. Moreover, injection of the CAT construct

with the similarly prepared liposomes resulted in lung expres-
sion of CAT, but failed to prevent the LPS-induced lung PMN
uptake. NIF expression in lungs also did not reduce the basal
level of PMN sequestration, a finding in agreement with the
lack of involvement of CD11/CD18 integrins in regulating the
normal marginated pool of PMN in the pulmonary vascular
bed (28).
We have shown previously that NIF gene transfer in cultured endothelial cells elicits NIF release into the culture conditioned medium (Malik, A.B., unpublished observation); however, we were not able to detect circulating NIF in mice (data not shown), which may reflect NIF’s binding to leukocytes (22, 23). In this regard, the localized release of NIF into the microcirculation by endothelial cells and other vessel wall cells and its rapid binding to β2 integrins on LPS-activated PMN may have the advantage of not interfering with the immune host-defense function of PMN. In contrast, the i.v. injection of anti-CD18 mAbs impairs PMN function and worsens infection (3, 5, 20, 34). The finding that NIF expression did not induce the leukocytosis that is characteristically observed after injection of anti-CD18 mAbs and in patients with leukocyte adhesion deficiency (LAD) (3) also suggests that the release of NIF by the transduced cells is a localized event. Therefore, in vivo transfer of the NIF gene can provide a means of continuously delivering NIF into the microcirculatory compartment to prevent PMN adhesion and migration of PMN across the vessel wall. Further specificity in NIF delivery may be effected through endothelial cell type–specific and antibiotic-inducible promoters.

In summary, we report that transfer of the NIF gene encodes the 41-kD antiadhesive protein in pulmonary microvesSEL wall cells of mice. NIF expression prevented LPS-induced lung PMN uptake and PMN migration into airspaces, and the associated lung vascular injury. We conclude that in vivo pulmonary vascular expression of NIF, a specific β2 integrin-binding protein, is a potentially useful gene transfer strategy in modulating the infiltration of PMN across the alveolar–capillary epithelial barriers and in preventing lung endothelial injury.

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References


In Vivo NIF Expression Prevents Lung Injury 2437