Interaction of a Specific Population of Human Embryonic Stem Cell–Derived Progenitor Cells with CD11b+ Cells Ameliorates Sepsis-Induced Lung Inflammatory Injury

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Human embryonic stem cells differentiated under mesoderm-inducing conditions have important therapeutic properties in sepsis-induced lung injury in mice. Single cell suspensions obtained from day 7 human embryoid bodies (d7EBs) injected i.v. 1 hour after cecal ligation and puncture significantly reduced lung inflammation and edema as well as production of tumor necrosis factor-α and interferon-γ in lungs compared with controls, whereas interleukin-10 production remained elevated. d7EB cell transplantation also reduced mortality to 50% from 90% in the control group. The protection was ascribed to d7EB cell interaction with lung resident CD11b+ cells, and was correlated with the ability of d7EB cells to reduce it also reduced production of proinflammatory cytokines by CD11b+ cells, and to endothelial NO synthase–derived NO by d7EB cells, leading to inhibition of inducible macrophage-type NO synthase activation in CD11b+ cells. The protective progenitor cells were positive for the endothelial and hematopoietic lineage marker angiotensin converting enzyme (ACE). Only the ACE+ fraction modulated the proinflammatory profile of CD11b+ cells and reduced mortality in septic mice. In contrast to the nonprotective ACE-cell fraction, the ACE+ cell fraction also produced NO. These findings suggest that an ACE+ subset of human embryonic stem cell–derived progenitor cells has a highly specialized anti-inflammatory function that ameliorates sepsis-induced lung inflammation and reduces mortality. (Am J Pathol 2011, 178:313–324; DOI: 10.1016/j.ajpath.2010.09.041)
strated marked reduction in mortality. Dampening of lung inflammation was the result of progenitor cells enriched with the endothelial and hematopoietic progenitor cell marker angiotensin-converting enzyme (ACE) and was largely ascribed to the interaction of these cells with CD11b+ cells in lungs. This interaction in turn mediated reduction in production of proinflammatory cytokines and high-output NO production by CD11b+ cells.

Materials and Methods

Differentiation of hESCs into Embryoid Bodies

hESCs (H1, XY, WiCell, and National Institutes of Health–approved WA01) were maintained on mitomycin-blocked mouse embryonic fibroblast feeders in hESC growth medium (Dulbecco’s modified Eagle’s medium and Ham nutrient mixture F-12) supplemented with 15% knockout serum replacement enriched with 4 ng/ml of human basic fibroblast growth factor-2, 1× nonessential amino acid, 1× glutamax-I, and 1× β-mercaptoethanol (all from Invitrogen Corp., Carlsbad, CA). Half of the medium was changed every 48 hours until the colonies were close to confluence. For differentiation induction, 2 to 2.5 × 10^5 hESCs were resuspended in 3 ml of stem cell medium (HEScGro; Millipore Corp., Billerica, MA) supplemented with 50 ng/ml of vascular endothelial growth factor and 50 ng/ml of bone morphogenetic protein-4, plated in one well of a six-well plate (Ultra-Low; Corning Inc., Corning, NY), and incubated at 37°C with 5% CO2. After 24 hours, 40 ng/ml of stem cell factor, 40 ng/ml of thrombopoietin, and 40 ng/ml of Fms-related tyrosine kinase-3 ligand (R&D Systems, Inc., Minneapolis, MN) were added to the cultures, followed by 25 ng/ml of granulocyte colony-stimulating factor, granulocyte-macrophage colony-stimulating factor, IL-6, and IL-3, and 3 U/ml of human erythropoietin at day 3½ of differentiation culture.

Differentiation of hESCs into Endothelial and Hematopoietic Progenitor Cells

d7EB cells were fractionated using fluorescein-activated cell sorting (FACS) for ACE and kinase insert domain receptor (KDR) expression. The isolated fractions were subcultured on fibronectin-coated plates in the presence of endothelial cell basal medium and 20 ng/ml of stem cell factor, 20 ng/ml of thrombopoietin, 20 ng/ml of Fms-related tyrosine kinase-3 ligand (Flt3) ligand (R&D Systems, Inc., Minneapolis, MN) were added to the cultures, followed by 25 ng/ml each of granulocyte colony-stimulating factor, granulocyte-macrophage colony-stimulating factor, IL-6, and IL-3, and 1 U/ml of human erythropoietin.

Lung Tissue Myeloperoxidase Content

Lung tissue neutrophil (polymorphonuclear leukocyte) uptake was assessed via determination of myeloperoxidase activity. Lungs were dried and homogenized in 1.0 ml of PBS (50 mmol/L, pH 6.0) with 5% hexadecyl-trimethylammonium bromide and 5 mmol/L of EDTA. Homogenates were sonicated, centrifuged at 4 × 10^6 g for 20 minutes and were frozen and thawed twice, followed by homogenization and centrifugation. The supernatant was mixed 1:30 (v/v) with assay buffer (0.2 mg/ml of o-dianisidine hydrochloride and 0.0005% H2O2), and absorbance change was measured at 460 nm for 3 minutes. Myeloperoxidase activity based on dry lung weight was calculated as the change in absorbance over time.

Lung Vascular Permeability Measurement

Lung capillary filtration coefficient was measured to quantify lung vascular permeability, as described previously. Venous outflow pressure was elevated using a computer-controlled two-way electronic pinch valve (P/N 98301-22; Cole-Parmer Instrument Co., Vernon Hills, IL) that channeled venous fluid into silicon tubing (1/16-inch i.d.). The weight change resulting from the venous pressure increase of 6 cm of water was recorded for 5 minutes. The weight recording has two exponential components that reflect the rapid expansion of vascular volume and the slower phase of transvascular fluid filtration. The amount of fluid filtered in 5 minutes was determined via logarithmic extrapolation of the slower component to time 0. The lung capillary filtration coefficient, a measure of lung vascular liquid permeability, was computed as milliliters per minute per centimeter of water per gram dry
weight by normalizing the estimated filtered fluid by time, venous pressure change, and lung dry weight.

**Isolation and Depletion of Lung CD11b+ Cells**

At 6 and 12 hours after CLP, lungs were removed from euthanized mice, minced into small pieces, and incubated in RPMI-1640 medium with 1% penicillin-streptomycin and 1% glucose for 30 minutes at 37°C and at 5% CO₂ in the presence of 1 mg/ml of collagenase type 1 and 50 U/ml of deoxyribonuclease I. After incubation, cells were filtered through a 70-µm cell strainer and washed with RPMI medium. The cells were incubated for 15 minutes at 4°C with CD11b magnetic beads (Miltenyi Biotek, Inc., Auburn, CA) and subsequently applied to MS columns (Miltenyi Biotek, Inc.) for their positive selection. Negatively selected cells were also collected.

For CD11b+ depletion studies, 24 hours before experimental procedures, mice were injected i.v. with gadolinium chloride, 10 µg/g body weight; anti–Gr-1, 50 µg per mouse; and anti–NK1.1, 50 µg per mouse. Depletion of CD11b+ cells was verified using flow cytometry.

**Histologic Analysis and Immunohistochemistry**

Mouse lungs were inflated with 10% formalin and embedded in paraffin. Formalin-fixed paraffin-embedded tissue samples were cut into 5-µm sections, mounted on slides (Starfrost Plus; Maenzel Glaeser, Braunschweig, Germany), and hydrated using an alcohol gradient. Slides were rinsed in distilled water, followed by antigen unmasking using a 10× concentrated retrieval solution (antigen decloaker solution; Biocare Medical Inc., Concord, CA) according to the manufacturer’s instructions, and rinsed in PBS for 5 minutes. For detection of human laminin A1, tissue sections were blocked with H₂O₂ rinsed in PBS for 5 minutes. For detection of human CA) according to the manufacturer’s instructions, and CD43 at 1:200 dilution in 2% normal donkey serum, 1% bovine serum albumin, and 0.1% Triton-X in PBS. Control cells were incubated with goat IgG, rabbit IgG, and mouse IgG, respectively, under the same conditions. After washing with PBS and repeat blocking for 1 hour, the cells were incubated with secondary antibodies Alexa 488 anti-goat IgG, Alexa 568 anti-rabbit IgG, or Alexa 568 anti-mouse IgG1 at 1:700 dilution for 1 hour at room temperature.

**Labeling for Stem Cell Tracking**

Single cell suspensions were created from d7EB cells via brief trypsinization. The cells were then fluorescently labeled via incubation with 10 µmol/L of carboxy-fluorescein diacetate (Green Tracker, C2925; Invitrogen Corp.) in serum-free medium for 30 minutes at 37°C. Labeling was confirmed at fluorescence microscopy, and cells were kept on ice until use. Carboxy-fluorescein diacetate-labeled green fluorescent cells, 5 × 10⁵, were injected i.v. through the facial vein. For cell tracking studies, mice were sacrificed at 6, 18, and 24 hours after injection, and 5-µm snap-frozen lung sections were cut. Tissues were counterstained with DAPI (4',6-diamino-2-phenylindole) and examined for native green fluorescence at confocal microscopy (Zeiss LSM 510 META confocal microscope; Carl Zeiss AG, Oberkochen, Germany). Sections were graded using a semiquantitative scale established by counting the number of fluorescent cells per 100 nuclei per power field examined at 40× magnification (0 = no florescence, 1 = <5 fluorescent cell, 2 = 1 to 5 fluorescent cells, 3 = >5 fluorescent cells). Ten power fields per slide were examined by two independent reviewers blinded to the animal group.

**Lung Edema Determination**

Extravascular lung water content was measured via determination of lung wet-to-dry weight ratios in which intravascular lung water was corrected using a hemoglobin assay of lung homogenates and peripheral blood.⁴

**Flow Cytometry**

The following antibodies were used for flow cytometry studies and FACS: mouse anti-CD11b Alexa Fluor 488–conjugated (clone M1/70; isotype, rat IgG₂a–488); mouse anti-F4/80 (clone 6F12; isotype, rat IgG₂a); mouse anti-NK1.1 PE-conjugated (clone PK136; isotype, mouse IgG₂a-PE); mouse anti-Gr1 PE-conjugated (clone RB6-8C5; isotype, mouse IgG₂a-PE); rat anti-mouse IFN-γ (clone XMG 1.2; isotype, rat IgG1); rat anti-mouse IL-10 (clone JES5-16E3; isotype, rat IgG₂b); biotin mouse anti-human TLR4 (clone HTA 125; isotype, mouse IgG₂a-biotin); and biotin anti-Annexin V (all from BD Biosciences Pharmingen, San Di-
ego, CA); mouse anti-KDR PE-conjugated (isotype, mouse IgG1, PE) and mouse anti-ACE fluorescein isothiocyanate-conjugated (isotype, goat IgG–fluorescein isothiocyanate) (both from R&D Systems, Inc.); rat anti-mouse TNF-α (clone MP6-XT22; isotype, rat IgG1; Invitrogen Corp.); rabbit polyclonal anti-NOS2 (clone N-20; isotype, goat IgG; Santa Cruz Biotechnology, Inc., Santa Cruz, CA); and NO-Cu-Fi, final concentration 10 μM/L (Intracellular Nitric Oxide Sensor Kit, catalog No. 96–0293; Strem Chemicals, Inc., Newburyport, MA). For intracellular staining, mouse lung cells were first fixed via incubation using 100 μL of fixation buffer (eBioscience, Inc., San Diego, CA) for 20 minutes at room temperature, and washed twice with 1 ml of permeabilization buffer (eBioscience, Inc., San Diego, CA). Consequently, 1 to 5 × 10^6 cells of antibodies against TNF-α, IL-10, IFN-γ, or inducible NO synthase (iNOS) were then added in 100-μL volume of permeabilization buffer, and cells were incubated for 30 minutes at 4°C. After an additional washing with 1 ml of permeabilization buffer, the cells were stained with secondary anti-rat–fluorescein isothiocyanate or anti-rabbit–PE (iNOS) conjugated antibody for 20 minutes at 4°C. The cells were resuspended in flow cytometry buffer and analyzed immediately. Intracellular NO staining with fluorescent-bound copper was performed according to the manufacturer’s instructions and in accordance with published protocols.17,18 Surface and intracellular marker expression were analyzed using software (LSR and CellQuest Pro; Becton Dickinson & Co., San Jose, CA).

**Enzyme-Linked Immunosorbent Assay**

Enzyme-linked immunosorbent assays were performed on supernatants of lung homogenates and cell culture supernatants using kits for mouse-specific TNF-α (Mouse TNF-α/TNFSF1A Quantikine ELISA Kit, MTA00); IFN-γ (Mouse IFN-γ Quantikine ELISA Kit, MIF00); and IL-10 (Mouse IL-10 Quantikine ELISA Kit, M1000) (all from R&D Systems, Inc.) according to the manufacturer’s instructions.

**Co-culture Experiments**

Single-cell suspensions of mouse lung CD11b+ and CD11b− cells were prepared and plated in 24-well plates at a concentration of 10^6 cells in 1 ml of complete RPMI-1640 for 1 hour. The supernatants were removed, and 2 × 10^5 d7EB cells in fresh medium were added either directly with the mouse cells or on the insert membrane of the Transwell system (HTS Transwell 0.4-μm pore size polycarbonate membrane; Corning, Inc.). As a control, CD11b+ cells were plated without d7EB cells. The co-cultured cells were or were not stimulated with 1 μg/ml of lipopolysaccharide (LPS) for 6 hours. At the end of the stimulation period, the supernatants were collected, and mouse-specific ELISAs (R&D Systems, Inc.) were performed on samples to detect the released mouse TNF-α, IFN-γ, and IL-10. This protocol was repeated for the experiments with ACE and KDR sorted d7EB cells.

**Assay for NO and NO-Derived Products**

Concentrations of NO and nitrate were determined using an NO analyzer (NOATM280; Sievers Instruments, Inc., Boulder, CO). NO concentrations were determined via analysis of nitrate accumulation in the culture supernatants of co-cultured d7EB cells and CD11b+ cells or CD11b− cells alone. Media aliquots were collected at the indicated times, and were injected directly into the analyzing cell containing the reducing solution (saturated sodium iodide in 50% acetic acid). The analysis was conducted at room temperature. Authentic NaNO₂ solutions of known concentrations were used as standards. For determination of NO production from human d7EB cells, cultures were washed twice with fresh serum-free media. Medium containing 1 μg/ml of LPS was replenished, and cells were undisturbed for the duration of the experiment. Media samples were collected at selected times, and NO production was assessed as nitrite accumulated in the media.

**NO Donor Experiments**

Single-cell suspensions of CD11b+ cells were prepared from septic mice as described above. The cells were plated in 24-well plates at a concentration of 10^6 cells in 1 ml of complete RPMI-1640 medium for 1 hour. The supernatants were removed, and fresh medium was added in the presence or absence of LPS, 100 ng/ml, plus a series of concentrations of the NO donor diethylenetriamine (DETA) NONOate for 12 hours. The range of concentrations was chosen to mimic approximately the amount of NO produced by d7EB cells as determined by direct measurements as described above. At the end of the experiment, supernatant free of cells was collected for cytokine measurements via ELISA.

**Generation of Conditioned Medium**

Human d7EB cells were plated in 24-well plates at concentration of 2 × 10^6 cells in 1 ml of growth medium. Cells were cultured overnight in the presence or absence of LPS, 1 μg/ml. The supernatant was collected, spun down for 10 minutes at 500g to remove possible cell contamination, and injected i.v. into septic mice 1 hour after CLP at a volume of 200 μL per mouse.

**Western Blot Analysis**

CD11b+ cells were either co-cultured with d7EB cells as described above, exposed to DETA NO, or cultured alone. After stimulation with LPS, 1 μg/ml, for 12 hours, the cells were lifted by gentle scraping. Human cells, which do not express CD11b, were removed via positive selection using CD11b magnetic beads (Miltenyi Biotek, Inc.). Live CD11b+ cells were lysed in 1× radioimmuno-precipitation assay lysis buffer containing protease inhibitor cocktail, 60 μL/10 ml PBS (Sigma-Aldrich Corp.). Lysates were centrifuged at 14,000 rpm for 10 minutes at 4°C. Supernatants were collected, and the protein concentration of each sample was measured using a bicinchoninic acid assay kit with bovine serum albumin as the
were observed for 6 months for development of teratomas. At the end of the observation period, the injection areas were dissected and stained with H&E. No evidence of teratomas was observed in any of the animals.

Statistical Analysis

Data were analyzed and figures generated using commercially available software (Prism version 4.03; GraphPad Software, Inc., San Diego, CA). Quantitative data are given as mean (SEM). Comparison between transplant and control groups was made using the nonparametric Mann-Whitney test. \( P < 0.05 \) was considered significant. Survival between intervention and control groups was compared using the log-rank test.

Results

d7EB Cell Transplantation Prevents Sepsis-Induced Lung Inflammatory Injury and Reduces Mortality in Mice

First, the effects of d7EB cells on sepsis-induced lung inflammatory injury and death induced via CLP were determined. Because the studies involved transplantation of human progenitor cells in mice, the question of xenotransplantation was considered. In mice treated with CsA, transplantation of human d7EB cells (500,000 cells in 200 \( \mu \)L of PBS) reduced mortality after lethal CLP, from 10% in control mice receiving mitomycin-blocked mouse embryonic fibroblasts (MEFs) to approximately 40% in the transplant group (Figure 1A). Although the CsA-immunosuppressed septic model proved useful in establishing a therapeutic role for human cells, there remained the important bias of pharmacologic immunosuppression in the sepsis model. Based on findings that ESCs and standard (Pierce Chemical Co., Rockford, IL). For each sample, 50 \( \mu \)g of protein was loaded onto lanes of NuPAGE 4% to 12% Bis-Tris gel (Invitrogen Corp.). Proteins were transferred to nitrocellulose membranes (Millipore Corp.). After incubation in blocking solution (5% dry milk in Tris-buffered saline solution with Tween 20) at room temperature for 1 hour, membranes were immunoblotted (24 hours at 4°C) with anti-iNOS rabbit polyclonal antibody (ab3523) and anti-eNOS mouse monoclonal antibody (ab76199) (both from Abcam Inc., Cambridge, MA), followed by secondary horseradish peroxidase-conjugated goat anti-rabbit or mouse (1:1000). Peroxidase labeling was detected using the ECL Western Blotting Detection System (GE Healthcare, Piscataway, NJ).

Tumorigenesis Assay

Five NOD/SCID mice were injected subcutaneously with 2 \( \times \) 10^6 d7EB cells in the dorsal region, and the animals

Figure 1. Transplantation of d7EB human progenitor cells improves sepsis-induced mortality. A: Transplantation of CsA-immunosuppressed mice with human d7EB cells (500,000 cells in 200 \( \mu \)L of PBS) 1 hour after abdominal sepsis induced by CLP significantly improved survival from 10% in the MEF-transplanted control group to 40%. \( N = 50 \) in each group, \( P < 0.0001 \), log-rank test. B: d7EB cell transplantation in immunologically intact mice significantly improved survival after CLP. d7EB and MEF control cells (500,000 in 200 \( \mu \)L of PBS) were injected i.v. 1 hour after CLP. At the end of 120 hours of observation, survival in the control group receiving MEF was 10% versus 50% in the d7EB cell recipient group. \( N = 50 \) in each group. \( P < 0.0001 \), log-rank test.

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Figure 2. Transplantation of d7EB human progenitor cells improves sepsis-induced lung inflammatory injury. A: Transplantation of d7EB cells reduced lung neutrophil sequestration, as measured using lung tissue myeloperoxidase activity, 24 hours after CLP. Controls were nonseptic mice. \( N = 15 \) in each group. \( P < 0.05 \); error bars represent SEM. B: d7EB cell transplantation reduced lung edema formation induced by CLP. Cells (500,000 in 200 \( \mu \)L of PBS) were injected 1 hour after CLP. Extravascular lung water content was measured 24 hours after CLP for determination of lung wet-to-dry-weight ratios. Controls are nonseptic mice. \( N = 15 \) in each group. \( P < 0.05 \); error bars represent SEM. C: Transplantation of d7EB cells dampened the increase in pulmonary vascular permeability, as measured using pulmonary capillary filtration coefficient (Kf,c), 24 hours after CLP. Controls were nonseptic mice. \( N = 5 \) in each group. \( P < 0.05 \); error bars represent SEM. D: Transplantation of d7EB cells improved lung architecture after CLP. Representative images of H&E-stained lung sections were obtained 24 hours after CLP. Magnification \( \times 20, \times 10 \). E: Reduced lung TNF-\( \alpha \) and IFN-\( \gamma \) production after d7EB cell transplantation. TNF-\( \alpha \), IFN-\( \gamma \), and IL-10 were measured in whole-lung tissue at the indicated times after induction of CLP. Transplantation of d7EB cells significantly reduced TNF-\( \alpha \) and IFN-\( \gamma \) production but did not change IL-10 production. \( N = 10 \) in each group. \( P < 0.05 \); error bars represent SEM.
adult bone marrow stromal cells\textsuperscript{19–22} might not be strongly immunogenic because they generally lack major histocompatibility complex II antigens;\textsuperscript{23} the effectiveness of these cells in immunocompetent septic mice without the complicating effects of previous immunosuppression was evaluated. Human d7EB cells in these mice were as protective against CLP-induced death as demonstrated in CsA-treated mice; that is, survival improved from less than 10\% at 48 hours after CLP in control mice to 50\% in the transplant group (Figure 1B). Also investigated were alterations in lung vascular permeability and edema formation at 24 hours after CLP. d7EB cell transplantation significantly reduced neutrophilic lung inflammation and lung edema and prevented lung endothelial barrier dysfunction in treated mice compared with the control group (Figure 2, A–D). The improvement in lung injury and survival was associated with decreased production of TNF-\(\alpha\) and IFN-\(\gamma\) in septic lungs, whereas there was no change in IL-10 production (Figure 2E).

Despite the absence of immunosuppression, d7EB cells were not immediately rejected, and their presence in recipient mouse lungs was verified up to 24 hours after cell transplantation. Increased numbers of cells were observed in the recipient lungs at 1 and 3 hours after transplantation, as demonstrated by human-specific laminin staining (Figure 3A). The number of cells in the lungs gradually decreased, but cells were still detectable during the first 24 hours (Figure 3B); however, no cells were visible at 48 hours after administration (data not shown). To determine whether the protection induced by d7EB cells could be ascribed to secreted factors, cultured d7EB cells were stimulated overnight with LPS, 1 \(\mu\)g/ml, or medium alone. Addition of LPS-conditioned medium obtained from d7EB cells did not alter mortality in mice that underwent CLP compared with controls injected with nonconditioned medium or PBS (Figure 3C), indicating that d7EB cells were essential for protection.

**Interaction of d7EB Cells and CD11b+ Cells Mediates Lung Protection**

Because the primary source of TNF-\(\alpha\) and IFN-\(\gamma\) in inflamed lungs is CD11b+ cells, that is, resident macrophages, blood monocytes, granulocytes, and natural killer cells,\textsuperscript{24} the possibility that d7EB cell interaction with host CD11b+ cells might induce an anti-inflammatory cytokine profile was investigated. CD11b+ cells were isolated from lungs of septic mice treated with either d7EB cells or MEFs. These cells were then stimulated with LPS in culture in the presence of the intracellular protein transport inhibitor Golgi stop. Cytokine staining showed that lung CD11b+ mouse cells obtained after d7EB cell transplantation produced significantly less TNF-\(\alpha\) and IFN-\(\gamma\) (Figure 4A) compared with lung CD11b+ cells from mice receiving control MEF. These responses were not observed in CD11b- cells (Figure 4A).

Whether d7EB transplantation alone influenced the CD11b+ cell population isolated from lungs of septic mice undergoing d7EB transplantation was addressed. No significant difference was observed in the number of monocytes (F4/80-positive), polymorphonuclear leukocytes (Gr-1–positive), and natural killer cells (NK1.1-positive) in the CD11b+ populations of the d7EB recipient and control lungs (Figure 4B).

To examine whether cell-cell interaction mediated the lung protection induced by d7EB cell transplantation, studies were performed in mice depleted of macrophage and monocyte populations using GdCl\(_3\) in combination with anti-Gr1 and anti-NK1.1 antibodies injected i.v. 24 hours before CLP challenge. Mortality was higher in these mice after CLP, which could not be prevented by d7EB cell transplantation (Figure 4C). Thus, d7EB cell interaction with macrophage or monocyte cells is important in the mechanism of protection.

To address whether direct interaction of mouse CD11b+ cells with d7EB cells was required for the shift to anti-inflammatory lung cytokine profile observed in mice in the cell transplantation group, lung CD11b+ cells were isolated from septic mice 12 hours after CLP and co-cultured with d7EB cells either directly or indirectly via separation using a 0.4-\(\mu\)m filter. Both direct and indirect co-cultures were stimulated with LPS for 6 hours, and the supernatants were analyzed for TNF-\(\alpha\) and IFN-\(\gamma\) production using a mouse-specific
ELISA. Only direct co-culture reduced TNF-α and IFN-γ production by the CD11b+ cells (Figure 4D).

Interaction of d7EB Cells with Septic Lung CD11b+ Cells Reduces NO Production

The observation that direct interaction of d7EB and CD11b+ cells was required for the anti-inflammatory phenotype of CD11b+ cells led to consideration of the involvement of NO, an important inflammatory mediator of sepsis. In the response. A time course study of NO production in cultured d7EBs over 6 hours demonstrated that d7EB cells constitutively produced NO and that they also strongly expressed eNOS (Figure 5, A and B).

Evidence suggests that NO supplementation in the form of NO donors exerts an anti-inflammatory effect in animal models of endotoxemia through inhibition of nuclear factor κB activation and iNOS expression, and decreased production of proinflammatory cytokines. Therefore, whether the NO-producing d7EB cells exerted a similar anti-inflammatory effect on CD11b+ cells was determined. d7EBs were co-cultured with CD11b+ cells, and nitrite concentration was measured in the supernatant after LPS stimulation. Supernatant NO concentrations at 6 and 12 hours after LPS stimulation of co-cultured d7EB and CD11b+ cells were significantly lower than the supernatant NO concentration from CD11b+ cells alone (Figure 5C). This effect could not be attributed to increased apoptosis because apoptosis was not significantly different in the co-cultured CD11b+ cells compared with the CD11b+ cells cultured alone (see Supplemental Figure S1 at http://ajpamjpathol.org). Analysis of iNOS expression in co-cultured CD11b+ cells showed decreased protein expression in LPS-stimulated CD11b+ cells interacting with d7EB cells (Figure 5D). This finding was supported by measurement of iNOS expression.
in CD11b+ cells isolated from the lungs of septic mice 12 hours after CLP, where decreased expression of the enzyme was again observed (Figure 5E). To further test the hypothesis that NO production by d7EB cells contributes to the anti-inflammatory phenotype of the mouse CD11b+ cells, an NO donor (DETA NONOate) was added directly to LPS-stimulated CD11b+ cells, this time without d7EB cells, at concentrations ranging from 0.5 to 2 μmol/L, and cytokine concentrations in the supernatant were measured. Reduction in TNF-α and IFN-γ production by the LPS-stimulated CD11b+ cells was observed (Figure 5F). In addition, iNOS expression in these cells after stimulation with LPS was reduced to levels comparable to those observed after co-stimulation with human d7EB cells (Figure 5G).

ACE Expression Defines the d7EB Cell Population Responsible for Dampening CD11b+ Cell Activation

ACE and KDR expression in d7EB cells was monitored because these markers are associated with differentiation of hematopoietic and endothelial progenitor cells.12,28 Neither ACE nor KDR were expressed in undifferentiated stem cells; however, increased cell surface expression of these markers was observed after onset of mesodermal differentiation, starting at day 3 and reaching maximum at day 7 to 8 when 23% of d7EB cells were ACE+KDR+ and 23% were ACE+KDR− (Figure 6A). To determine whether these two markers identified distinct
lineages of mesodermally differentiated cells, FACS-fractionated d7EB cells were subcultured based on their expression of either ACE or KDR. After 4 days in subculture, ACE+KDR+ cells gave rise to endothelial cell colonies with a characteristic architecture and positive staining for VE-cadherin and von Willebrand factor. When these cells were cultured in a two-dimensional Matrigel system (BD Biosciences Pharmingen), they gave rise to colonies with a characteristic architecture and positive staining for VE-cadherin and von Willebrand factor. When these cells were cultured in a two-dimensional Matrigel system (BD Biosciences Pharmingen), they gave rise to
were fractionated and used for survival studies compar-
only by the ACE-expressing cells, both ACE
ACE and KDR with NO production. NO was produced
differential NO production by the ACE
were not protective (Figure 6F). To examine whether the
ACE/H11002/KDR, whereas ACE/H11001/KDR
reproduced the protective effect of the mixed d7EB pop-
ulation in survival against sepsis, whereas ACE
ACE/KDR, whereas ACE/KDR− fraction showed great efficiency in pro-
duction of hematopoietic colonies (Figure 6C). To deter-
mine the importance of ACE and KDR markers in the
observed protective phenotype against sepsis, fraction-
ated d7EB cells were co-cultured according to their ex-
pression of ACE and KDR with the CD11b+ cells from
septic mice as described above. The co-cultured cell
fractions were stimulated in vitro with LPS for 12 hours,
and mouse TNF-α and IFN-γ production was monitored.
Both the ACE+/KDR− fraction and the ACE+/KDR+ fraction significantly reduced TNF-α and IFN-γ production (Figure 6D), indicating the critical role of the progen-
itor cells expressing only ACE in reducing TNF-α and
IFN-γ production by CD11b+ cells. In addition, a strong
cell surface co-expression of ACE and toll-like receptor 4
(TLR4) was observed in d7EB cells, suggesting an asso-
ciation between ACE expression and the capability to
respond to LPS (Figure 6E).

To further validate these observations, the ACE+ cells
were fractionated and used for survival studies compar-
ing them with the ACE− fraction. Only ACE+ d7EB cells
reproduced the protective effect of the mixed d7EB pop-
ulation in survival against sepsis, whereas ACE− cells
were not protective (Figure 6F). To examine whether the
reduced production of cytokines was associated with
differential NO production by the ACE+ progenitor cells,
flow cytometry was used to determine co-expression of
ACE and KDR with NO production. NO was produced
only by the ACE-expressing cells, both ACE+/KDR+ and
ACE+/KDR−, whereas ACE− cells did not produce NO
(Figure 6G).

Discussion

hESCs, derived from the inner cell mass of the pre-im-
plantation blastocyst, are defined by their ability to self-
renew and to differentiate into all types of mature
cells.29,30 Although these cells and their derivatives in
different stages of differentiation have been used for cell
therapy applications in animal models of cardiovascular
disease,31–33 peripheral vascular disease,12,34 and cen-
tral nervous system disorders,21,22,35,36 their potential in
preventing sepsis-induced lung inflammatory injury char-
acteristic of adult respiratory distress syndrome has not
been addressed. The present study demonstrates for the
first time the role of a population of progenitor cells de-
rivied from hESCs in preventing lung inflammatory injury
induced by sepsis and improving survival in mice. The
results show that the protection is the result of the subset
of cells that are ACE+ to respond to LPS by producing
eNOS-derived NO. These cells functioned by moderating
the pro-inflammatory cytokine production of the host im-
mune CD11b+ cells and reducing the high output of
iNOS-derived NO cells and thereby mitigating lung in-
flammatory injury.

The protective effects of hESC-derived progenitor cells
were associated with decreased production of the pro-
inflammatory cytokines TNF-α and IFN-γ and mainte-
nance of the production of the major anti-inflammatory
cytokine IL-10. It was demonstrated that protection by
hESC-derived progenitor cells was the result of the inter-
action of these cells with resident lung CD11b+ cells.
Direct interaction of hESC-derived progenitor cells with
CD11b+ cells was required to create the anti-inflamma-
tory environment in lungs because neither injection of
septic mice with d7EB-conditioned medium nor indirect
cocultures reproduced the salutary changes in the cy-
tokine profile.

Because of the need for cell-cell interaction, the pos-
sibility was considered that paracrine factors with suffi-
cient diffusing capacity and instability in solution such as
NO might be involved. d7EB progenitor cells constitut-
ively produced eNOS-derived NO in amounts compara-
tible to those of immune cells, and strongly expressed
eNOS. Co-culturing CD11b+ cells with d7EB cells signifi-

cantly reduced NO production in the culture supernatant
after stimulation with LPS. This reduction in NO produc-
tion was coupled with inhibition of iNOS expression in the
cocultured CD11b+ cells and in CD11b+ cells isolated
from lungs of septic mice that received transplanted
d7EB cells.

eNOS-derived NO is beneficial in maintaining vascular
endothelial integrity,38 and eNOS-derived NO production
suppresses nuclear factor κB activity, decreases the
transcription of iNOS and intercellular adhesion mole-
cule-1, and prevents lung injury and death due to en-
do
toxin.39 Thus, NO produced by d7EB cells may protect
against iNOS activation and resultant high NO produc-
tion, which has known deleterious effects on the host.40
To test this hypothesis, CD11b+ cells were exposed to
an NO donor at a concentration range that induces NO
release equivalent to the amount of NO produced by
d7EB cells. This experiment reproduced the decreased
generation of pro-inflammatory cytokines and reduced
the iNOS expression observed in the CD11b+ cells co-
cultured with human d7EB cells. Thus, the results sug-
gest a critical role of eNOS-derived NO by d7EB cells in
down-regulating iNOS activation in CD11b+ cells and,
thereby, dampening the production of pro-inflammatory
cytokines.

These studies provide a mechanistic basis by which
d7EB cells prevent sepsis-induced lung inflammatory in-
jury. hESC-derived d7EB cells responsible for the protec-
tive phenotype were identified and found to be enriched
in two cell surface markers, ACE and KDR. It was also
demonstrated that the population consisting of
ACE+/KDR+ and ACE+/KDR− progenitor cells gave rise
in culture to endothelial and hematopoietic colonies, re-
spectively. In addition, the ACE-expressing cells ex-
pressed TLR4, the receptor sensing LPS, and produced
NO. In contrast, ACE− cells did not exhibit these char-
acteristics. The functional importance of these observations was reinforced by the ability of ACE+ fractions of d7EB cells to modulate the inflammatory cytokine production profile of CD11b+ host cells through direct cell-cell interaction. ACE is constitutively expressed on the surface of endothelial and hematopoietic precursors, and ACE has been identified as a marker of hematopoietic stem cells present at all stages in the ontogeny of the human hematopoietic system and as a novel marker of hemangioblasts differentiating from hESCs. Evidence also points to an important role of ACE in sepsis. ACE knockout mice exhibited improved lung injury scores after acid aspiration, endotoxin challenge, and peritoneal sepsis. Cohort studies in humans have shown a correlation between ACE polymorphisms and susceptibility to and death from adult respiratory distress syndrome, and decreased plasma concentrations of ACE have been observed in patients with adult respiratory distress syndrome and sepsis. A close homologue of ACE, ACE 2, has been identified as a key factor in protection from adult respiratory distress syndrome, and it also functions as a critical in vivo receptor for severe acute respiratory syndrome. A limitation of the present study is that the immunologic properties of human progenitor cells were studied in a xenograft model, introducing biases related to cross-species barriers. However, there is no reason to believe that species incompatibility affects the validity of the proposed mechanism, which involves interaction with host CD11b+ cells and modulation of NO production. Human cells were not acutely eliminated by the immune-competent mice because cells were detected in lungs for 24 hours after injection. These data are in accord with observations in syngeneic mouse mesenchymal stem cells and may be related to the immunosuppressive nature of the human cells. It is likely, however, that some degree of immunosuppression will be required for longer term xenotransplantation studies.

Although, to our knowledge, the present study is the first to describe the function of mesodermally differentiated hESCs in a model of lethal sepsis and lung injury, other studies have used bone marrow mesenchymal stromal cells in similar models. These studies demonstrated an anti-inflammatory benefit of these cells but showed variable effects on the production of anti-inflammatory and pro-inflammatory cytokines. In a polymicrobial sepsis model, the beneficial effect of bone marrow stem cells was attributed to stem cell–induced production of IL-10 by host macrophages, spingosine-1-phosphate production by stem cells, and homing of stem cells to lungs through integrin expression. hESC-derived progenitor cells represent a much earlier developmental stage than bone marrow mesenchymal stem cells, and they have demonstrated ability to differentiate into niche-specific mature cells, a characteristic that distinguishes them from adult mesenchymal stromal cells. Although the importance of the interaction of hESC-derived cells with host CD11b+ cells and the association of ACE expression with the protective phenotype were observed, factors such as homing to a niche and the engrafment potential of hESC-derived cells that may also be important in the observed protection cannot be ruled out. Findings of the present study demonstrate that d7EB cells, a population of hESC-derived progenitor cells, constitute a novel immunomodulatory cell population with therapeutic potential in sepsis that may have clinical application in cell-based therapy.

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