Pulmonary Epithelial Cell Expression of GM-CSF Corrects the Alveolar Proteinosis in GM-CSF–deficient Mice

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Abstract

Mutation of the granulocyte-macrophage colony-stimulating factor (GM-CSF) gene by homologous recombination caused alveolar proteinosis in mice. To further discern the role of GM-CSF in surfactant homeostasis, the synthesis of GM-CSF was directed to the respiratory epithelium of GM-CSF-null mutant mice (GM−/−) with a chimeric gene expressing GM-CSF under the control of the promoter from the human surfactant protein-C (SP-C) gene. Transgenic mice bearing the SP-C-GM-CSF construct (SP-C-GM+) were bred to GM−/− mice resulting in complete correction of alveolar proteinosis in bitransgenic GM−/−, SP-C-GM+ mice. No effects of the transgene were found outside the lung. GM-CSF was increased in bronchoalveolar lavage fluid of the bitransgenic mice. Surfactant proteins-A and -B and phospholipid in bronchoalveolar lavage fluid were normalized in the GM−/−, SP-C-GM+ mice. SP-A, -B, and -C mRNAs were unaltered in lungs from GM-CSF–deficient and –replete mice. Expression of GM-CSF in respiratory epithelial cells of transgenic mice restores surfactant homeostasis in GM−/− mice. From these findings, we conclude that GM-CSF regulates the clearance or catabolism rather than synthesis of surfactant proteins and lipids. (J. Clin. Invest. 1996. 97:649–655.) Key words: alveolar proteinosis • transgenic mice • granulocyte-macrophage colony-stimulating factor (GM-CSF) • surfactant homeostasis • gene correction

Introduction

Granulocyte-macrophage colony-stimulating factor (GM-CSF) is a 23-kD glycoprotein (1) encoded by a single gene containing four exons spanning 2.4 kb of the mouse genome (2). While GM-CSF was initially isolated from conditioned media from pulmonary cells, GM-CSF is expressed in a variety of hematopoietic cell types and nonhematopoietic cells and cell lines (3–8). GM-CSF has potent hematopoietic regulatory properties and was named for its ability to stimulate prolifera-
Methods

Construction of the SP-C-GM-CSF chimeric gene. Cloning vector p3.7-tpA, comprised of the human SP-C promoter region, nucleotides −3683 to +18 (3.7SP-C) (21, 22), followed by a multiple cloning site, SV40 small t-intron, and polyadenylation signal, was used to construct the chimeric gene in a pUC18 plasmid, as described previously (23). The coding region of mouse GM-CSF cDNA, bp 174–620, was isolated from pMFGmuGM-CSF (10). Ends were modified with addition of EcoRI linkers and inserted into the EcoRI site of p3.7-tpA to generate p3.7GM-tpA.

Production of transgenic mice. The SP-C-GM-CSF chimeric gene was excised from the parental plasmid p3.7GM-tpA by NdeI/NcoI digestion and purified by agarose gel electrophoresis in the absence of ethidium bromide. The DNA was recovered using Qiaex Gel extraction kit (QIAGEN Inc., Chatsworth, CA) and dialyzed in 5 mM Tris-HCl, pH 7.5, and 0.1 mM EDTA for 36 h. Donor eggs from FVB/N mice (GM−/−) were prepared for pronuclear injection. Sperm was obtained from GM-CSF-null mutant mice (GM−/−), described previously (16). Tail DNA was digested with BamHI and Southern blotted, using the EcoRI fragment of GM-CSF cDNA from p3.7GM-tpA to probe for wild-type (9.2 kb), null mutant (4.1 kb), and SP-C-GM-CSF-transgenic (0.5 kb) alleles. SP-C-GM-CSF transgenic founders were back-crossed with GM−/− mice to generate pups that were homozygous for wild-type (9.2 kb), GM−/−, GM−/+ mice, and GM+/− littermates were used as age-matched controls as indicated.

Processing and staining of tissues for histopathology. Lungs were inflation fixed with 4% paraformaldehyde/phosphate-buffered saline (PBS), pH 7.4, for 24 h, as described previously (24). Tissue was then washed in PBS, dehydrated in a series of alcohols, and embedded in paraffin. Hematoxylin and eosin staining was used for histological analysis of paraffin sections.

Immunohistochemistry. Paraffin sections of 5 μm were used to stain for surfactant protein. Anti–SP-B (28031) is a rabbit anti–bovine polyclonal antibody that recognizes mature SP-B (25). Anti–SP-B was detected using a secondary antibody and detection system (Vectastain ABC anti–rabbit immunohistochemical horseradish peroxidase kit; Vector Labs, Inc., Burlingame, CA). Tissues were counterstained with hematoxylin and eosin.

RNA isolation. RNA was isolated by a modification of the guanidium-thiocyanate method described by Chomczynski and Sacchi (26). Briefly, tissues were homogenized in guanidinium-thiocyanate solution kit (QIAGEN Inc., Chatsworth, CA) and dialyzed in 5 mM Tris-HCl, pH 7.5, and 0.1 mM EDTA for 36 h. Donor eggs from FVB/N mice (GM−/−) were prepared for pronuclear injection. Sperm was obtained from GM-CSF-null mutant mice (GM−/−), described previously (16). Tail DNA was digested with BamHI and Southern blotted, using the EcoRI fragment of GM-CSF cDNA from p3.7GM-tpA to probe for wild-type (9.2 kb), null mutant (4.1 kb), and SP-C-GM-CSF-transgenic (0.5 kb) alleles. SP-C-GM-CSF transgenic founders were back-crossed with GM−/− mice to generate pups that were homozygous for wild-type (9.2 kb), GM−/−, GM−/+ mice, and GM+/− littermates were used as age-matched controls as indicated.

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Figure 1. Gene constructs. (A) Wild-type mouse GM-CSF gene locus. (B) GM-CSF-null allele produced by gene targeting with the neomycin gene replacing exons 3 and 4, intronic sequences, and 3′ UTR of the mouse GM-CSF gene. (C) SP-C-GM-CSF chimeric gene containing the human SP-C gene promoter (3.7SP-C), coding sequence of mouse GM-CSF cDNA, SV40 small t-intron, and polyadenylation signal. BamHI digestion of genomic DNA yields diagnostic fragments for wild-type (9.2 kb), GM−/− (4.1 kb), or SP-C-GM-CSF transgene (0.5 kb) as indicated. B, BamHI; X, XbaI; H, HindIII; E, EcoRI; Neo, neomycin.
pg/ml were diluted and repeated. To assay saturated phosphatidylcholine (Sat PC) in BAL fluid, mouse lungs were lavaged and Sat PC was determined as described previously (27a).

**Results**

Promoter sequences (3.7SP-C) from the human SP-C gene, nucleotides −3683 to +18, were used to construct a chimeric gene directing expression of mouse GM-CSF cDNA in the respiratory lung epithelium of wild-type and GM−/− mice (Fig. 1, A–C). Ova from GM+/+ females were fertilized with sperm from GM−/− males. At the pronuclear stage of development, the SP-C-GM-CSF chimeric gene was injected into the ova, which were transferred to pseudopregnant foster mothers. Founders were screened by Southern blotting of tail DNA digested with BamHI, using a mouse cDNA probe (Fig. 2). Genotypes GM−/−, SP-C-GM− or GM+/−, SP-C-GM+ were determined by the presence of 9.2-, 4.1-, or 0.5-kb bands for

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White blood cell differentials were performed on peripheral blood smears from GM−/−, SP-C-GM+; GM−/−, SP-C-GM−; and GM+/−, SP-C-GM− animals. Results show percentage of monocytes, lymphocytes, and granulocytes counted; mean±SD; n = 5 individuals of each genotype.

![Figure 2. Southern blot analysis of GM−/−, SP-C-GM+ mice. Genotype was determined by Southern blotting of BamHI-digested tail DNA, using the mouse GM-CSF cDNA probe. Fragment sizes are indicated on the left in kilobases. Lanes 1 and 2 contain DNA from line 48, GM−/−, and GM+/−, respectively. Lanes 3 and 4 represent DNA from line 48, GM+/−, SP-C-GM+; and GM−/−, SP-C-GM+, respectively. Lane 5 is representative of line 59 GM−/−, SP-C-GM+ animals.]

![Figure 3. Lung histology in GM-CSF–deficient and –replete mice. Lungs from adult mice were inflation fixed and stained with hematoxylin and eosin. Alveolar proteinosis, enlarged foamy macrophages, and lymphocytic infiltration were consistent findings in GM−/− mice (A). Alveolar proteinosis was corrected in GM−/−, SP-C-GM+ mice from line 48 (B) and line 59 (C) in association with increased numbers of lung macrophages. Marked alveolar infiltration with macrophages was noted in the F16, GM+/−, SP-C-GM+ mouse (D). At least four individual mice were assessed from lines 48 and 59, ×340.]

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Three distinct GM/+−, SP-C-GM + transgenic mice founders were backcrossed to GM−/− mice to obtain GM−/−, SP-C-GM+ pups. One founder mouse (F16) failed to breed and was moribund at 4 mo; histology of lungs from this animal demonstrated marked pulmonary infiltration with alveolar macrophages and no alveolar proteinosis. Two other GM+/−, SP-C-GM+ mice, designated F48 and F59, were used to establish separate lines. Genotypes of offspring were as expected from Mendelian inheritance, indicating that the transgene had inserted into an autosomal locus in each line. Litters were of expected sizes, and pups were healthy with normal growth and development. Mice from each line have been bred to the F4 generation and are now 11 mo old and are apparently healthy.

There were no apparent systemic effects of GM-CSF, and effects of the transgene were restricted to the lung. Peripheral blood counts were similar in GM−/−, SP-C-GM+; GM−/−, SP-C-GM−; and GM+/−, SP-C-GM− mice (Table I). Likewise, cell counts of peritoneal washes from GM−/−, SP-C-GM+; GM−/−, SP-C-GM−; and GM+/−, SP-C-GM− mice were indistinguishable, 5.0±1.8 (n = 3), 4.4±2.4 (n = 5), and 3.3±0.3 (n = 4) × 10⁶ cells recovered mean±SD, respectively. Values for peripheral blood and peritoneal wash cell counts were within ranges established in the literature (28, 29). Histopathologies of spleens from wild-type, GM−−−, and bitransgenic mice were identical. The numbers of MAC-1–reactive cells were increased in BAL fluid from the GM−−−, SP-C-GM+ mice compared with wild-type and GM−−− mice (lacking the SP-C-GM transgene) as assessed by flow cytometry, a finding consistent with the increased numbers of alveolar macrophages observed histologically (Fig. 3, A–D). Numbers of cells collected by BAL were increased similarly in GM+/−, SP-C-GM+ and GM−−−, SP-C-GM+ lines and were not different in mice from lines 48 or 59. BAL cell counts were 0.074±0.02 × 10⁶ mean±SD (n = 4) in GM+/− mice compared with 0.72±0.3 × 10⁶ (n = 4) in line 48 and 0.79±0.32 × 10⁶ (n = 4) in line 59 of the GM+/−, SP-C-GM+ mice. Similarly, cell numbers were 0.47±0.1 × 10⁶ (line 48) and 1.1±0.7 (line 59) n = 2 each in the GM−−−, SP-C-GM+ mice.

Proteinaceous accumulations in the alveolar spaces were not detected in lungs from GM−−−, SP-C-GM+ and GM+/−, SP-C-GM− (Fig. 4, C and E), or GM+/+ mice (16), in sharp contrast to lungs from GM−−−, SP-C-GM− littermates, wherein eosinophilic material filled the alveolar spaces in all lobes of the lungs, that was intensely stained by SP-B antibody (Fig. 4, A and B). The intensity of anti–SP-B staining of type II cells in GM−−−, SP-C-GM+ and GM+/−, SP-C-GM− mice (Fig. 4, D and F) was similar to that of GM+/+ wild-type mice.

Figure 4. SP-B in lungs of GM−−− and GM+/−, SP-C-GM+ mice. Lungs were inflation fixed and stained with hematoxylin and eosin (A, C, and E) or stained for SP-B (B, D, and F). Immunohistochemical staining with anti–SP-B antibody selectively stained type II cells in lung sections from GM−−−, SP-C-GM+ mice, but did not stain alveolar material (D and F). Intense staining of alveolar material was noted in the GM−−−, SP-C-GM− lung (B), corresponding to the proteinaceous infiltrates noted by hematoxylin and eosin staining (A). Lungs from GM+/−, SP-C-GM− mice stained with hematoxylin and eosin (E) and anti–SP-B (F) were indistinguishable from those of wild-type GM+/+ animals (not shown). Sections are representative of offspring from two separate founder SP-C-GM-CSF lines, assessing n = 3 mice per line.
SP-A, SP-B, and GM-CSF were measured in BAL fluid from GM−/−, SP-C-GM−; GM−/−, SP-C-GM+; and GM+/+, SP-C-GM− mice (Fig. 5). SP-A and SP-B levels in GM−/−, SP-C-GM− mice were ≥ 10-fold higher than in GM+/+, SP-C-GM− mice. The levels of SP-A and SP-B in GM+/+, SP-C-GM− mice did not differ significantly from GM+/+ mice (data not shown). SP-A and SP-B in BAL collected from GM−/−, SP-C-GM+ mice were not different from those in GM+/−, SP-C-GM− and GM+/+ mice. GM-CSF concentrations were below the level of detection in BAL from either GM−/−, SP-C-GM− or GM+/+, SP-C-GM− mice. GM-CSF in BAL fluid from GM−/−, SP-C-GM+ mice ranged from 151 to 438 pg/ml in both mouse lines. GM-CSF was below the level of detection in the serum from all mice, including the SP-C-GM+ transgenic mice (data not shown).

SP-A, SP-B, and SP-C mRNAs in GM−/−, SP-C-GM−; GM+/−, SP-C-GM−; and GM−/−, SP-C-GM+ were not different, suggesting that the correction of alveolar proteinosis in GM−/−, SP-C-GM+ mice was not directly related to changes in expression of surfactant protein mRNAs (Fig. 6).

Sat PC content in BAL from GM−/−, SP-C-GM− mice was ≥ 10-fold higher than that in BAL from GM+/−, SP-C-GM− mice (Fig. 7). Sat PC content in GM−/−, SP-C-GM+ mice did not differ significantly from GM+/−, SP-C-GM− or GM+/+ mice, demonstrating that expression of GM-CSF in the lungs of GM−/− mice normalized alveolar surfactant phospholipids and proteins.

**Discussion**

The local expression of murine GM-CSF in respiratory epithelial cells of lungs of transgenic mice corrected the alveolar proteinosis and lymphocytic infiltrates characteristic of the lungs of GM−/− mutant mice. The accumulation of surfactant proteins and phospholipids was resolved in the absence of detectable systemic effects of the transgene, supporting the hypothesis that the local production of GM-CSF by lung cells influenced surfactant homeostasis mediated by respiratory epithelial cells and alveolar macrophages.

The marked accumulations of surfactant in GM−/− mice and the resolution of the alveolar proteinosis in the GM−/−, SP-C-GM− mice were not observed in the “corrected” bitransgenic mice (Fig. 4, C and D).

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![Figure 5](http://www.jci.org)  
**Figure 5.** GM-CSF and surfactant proteins in GM−/− and GM+/−, SP-C-GM− mice. SP-A, SP-B, and GM-CSF were measured in BAL fluid from GM−/−, SP-C-GM−; GM−/−, SP-C-GM+; and GM+/+, SP-C-GM− mice (n = 4). SP-A and SP-B levels in GM−/−, SP-C-GM− mice were ≥ 10-fold higher than in GM+/−, SP-C-GM− mice; the latter were also within the range of healthy GM+/+ mice (data not shown). GM-CSF was not detectable in BAL from either GM−/−, SP-C-GM− or GM+/+, SP-C-GM− mice. SP-A and SP-B in BAL collected from GM−/−, SP-C-GM− and GM+/+ mice. Values are mean±SD, n = 4 per group. ND, not detectable.

![Figure 6](http://www.jci.org)  
**Figure 6.** S1-Nuclease assay for SP-A, -B, and -C mRNA. SP-A, SP-B, and SP-C mRNAs were assessed in lungs from GM−/−, SP-C-GM−; GM−/−, SP-C-GM−; and GM−/−, SP-C-GM+ mice. Sizes of protected fragments are indicated on the right in nucleotides (nt). When normalized for loading using ribosomal L32 surfactant protein mRNA were not different in the various lines tested. There was no statistically significant difference found between the genotypic groups. n = 4 by ANOVA.
SP-C-GM+ mice occurred independently of changes in surfactant protein mRNAs or intracellular SP-B staining. These findings suggest that altered surfactant metabolism, reuptake, or catabolism, rather than increased production of surfactant components, are involved in the PAP in GM−/− mice. The recent findings that surfactant phospholipid and 125I-SP-A clearance was markedly decreased in the GM−/− mice and that rates of phospholipid synthesis and secretion were relatively unchanged provide further support for the role of GM-CSF in surfactant clearance pathways (27a). On the other hand, surfactant synthesis and surfactant protein RNA content were not altered in the GM−/− mice in spite of the marked accumulation of surfactant proteins and lipids. Thus, counterregulatory mechanisms failed to decrease steady state surfactant synthesis that might have ameliorated abnormalities of surfactant accumulation in the PAP mice. These findings may reflect disordered counterregulatory mechanisms or that steady state surfactant concentrations are maintained primarily by the regulation of recycling and catabolism rather than by pathways that modulate surfactant biosynthesis.

In the present study, the human SP-C promoter (3.7SP-C) was used to direct synthesis of GM-CSF to alveolar type II cells and to distal bronchiolar epithelial cells in the lung. The h3.7SP-C promoter has been used extensively to express genes in a cell-specific manner in the lungs of transgenic mice. Expression of the exogenous cDNAs or mini-genes is consistently restricted to type II epithelial cells and distal bronchiolar epithelial cells (for review see reference 30). We were unable to detect SP-C-GM-CSF mRNA by either RT-PCR or Northern blot analysis, even though GM-CSF protein was readily detected in the BAL fluid of the SP-C-GM+ mice. The endogenous GM-CSF mRNA is frequently undetectable in biological samples, likely due to its short biological half-life (3). Instability of the GM-CSF mRNA may account for difficulty in detecting the transgenic GM-CSF mRNA. Nevertheless, restriction of pathological findings to the lung, increased concentrations of GM-CSF in the BAL fluid, and absence of extrapulmonary effects seen in the GM−/−, SP-C-GM+ mice suggest that the SP-C-GM-CSF mRNA is produced in the lung and is translated into biologically active GM-CSF. The finding that GM-CSF mRNA was not readily detected in the transgenic mice might be related to loss of transgenic mice expressing high levels of GM-CSF mRNA, although we did not observe increased perinatal death of SP-C-GM founder mice.

GM-CSF was readily detected in BAL but not in peripheral blood from the bitransgenic mice, providing further support for the concept that GM-CSF is acting locally, rather than peripherally, to correct the alveolar proteinosis in the GM−/− model. Peripheral blood leukocytes, splenic histology, and the lack of changes in peritoneal macrophages support the likelihood that the effects of the SP-C-GM-CSF transgene were not mediated by extrapulmonary GM-CSF. In contrast, transgenic mice expressing GM-CSF under control of the Moloney murine leukemia virus promoter developed abnormalities of tissue macrophage cell populations that were not observed in the present studies (8). It remains possible, however, that the local production of GM-CSF in the lung influences cells of granulocytic and monocytic cell lineages at distant sites and that the production of GM-CSF in the lung has provided signals necessary for appropriate granulocytic-monocytic maturation in other organs that corrects the abnormalities of surfactant homeostasis seen in the lung.

While alveolar macrophage numbers were not substantially altered in the lungs of GM−/− mice (16), macrophages became increasingly foamy and laden with lipids and surfactant proteins. Marked perivascular and peribronchiolar lymphocytic infiltration were also noted in the lungs of GM−/− mice. Both of these histologic features of the GM−/− mice were resolved in the bitransgenic mice expressing GM-CSF in the lung. The mechanisms involved in the resolution of PAP and restoration of macrophage morphology have not been clarified in the present studies. While morphology of the macrophages appears to be normal in the bitransgenic mice, the numbers of alveolar macrophages isolated in the lungs by alveolar lavage were increased in the GM−/−, SP-C-GM+ mice. These increased numbers of alveolar macrophages may contribute to more rapid clearance or catabolism in the corrected bitransgenic mice. Immunohistochemical staining for proliferating cell nuclear antigen was observed in a small percentage of macrophages in both wild-type and corrected mice, suggesting that local proliferation of macrophages may occur in the lungs of the GM−/− mice (data not shown). Since GM-CSF exerts proliferative, chemoattractant, and activating effects on alveo-
lar macrophages, the local proliferation of macrophages may not be the only mechanism resulting in the increased numbers and activity of alveolar macrophages in the corrected transgenic mice.

The present findings support a model by which local production of GM-CSF in the lung corrects PAP in the GM−/− mouse. However, the sites of expression and regulatory role of GM-CSF in normal lung physiology have not been clarified. While rat type II epithelial cells express GM-CSF in vitro (7), the normal in vivo sites and levels of GM-CSF protein or receptors in the lung remain to be elucidated. The relative instabilities of GM-CSF mRNA and protein complicate the analysis of potential signaling pathways that might be involved in regulation of surfactant homeostasis by GM-CSF and its receptors in the normal lung in vivo. It is therefore unclear whether surfactant homeostasis in the wild-type mouse and the abnormalities of surfactant homeostasis seen in GM-CSF mouse are mediated by interactions of GM-CSF with alveolar macrophages, type II epithelial cells, or both cell types.

Pulmonary alveolar proteinosis is a relatively uncommon, clinical syndrome with pathological features of surfactant lipid and protein accumulation similar to the findings of the GM−/− mouse. Pulmonary alveolar proteinosis has been associated with metabolic disease and acquired disorders of the hematopoietic systems (31). The present findings that the local production of GM-CSF corrects abnormalities of surfactant homeostasis in the bitransgenic GM−/−, SP-C-GM+ mice support the important role of GM-CSF in surfactant homeostasis and the hypothesis that abnormalities of GM-CSF or its receptors may be involved in the pathogenesis of PAP in humans.

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