Familial pulmonary alveolar proteinosis caused by mutations in CSF2RA

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Primary pulmonary alveolar proteinosis (PAP) is a rare syndrome characterized by accumulation of surfactant in the lungs that is presumed to be mediated by disruption of granulocyte/macrophage colony-stimulating factor (GM-CSF) signaling based on studies in genetically modified mice. The effects of GM-CSF are mediated by heterologous receptors composed of GM-CSF binding (GM-CSF-Rα) and nonbinding affinity-enhancing (GM-CSF-Rβ) subunits. We describe PAP, failure to thrive, and increased GM-CSF levels in two sisters aged 6 and 8 yr with abnormalities of both GM-CSF-Rα–encoding alleles (CSF2RA). One was a 1.6-Mb deletion in the pseudoautosomal region of one maternal X chromosome encompassing CSF2RA. The other, a point mutation in the paternal X chromosome allele encoding a G174R substitution, altered an N-linked glycosylation site within the cytokine binding domain and glycosylation of GM-CSF–Rα, severely reducing GM-CSF binding, receptor signaling, and GM-CSF–dependent functions in primary myeloid cells. Transfection of cloned cDNAs faithfully reproduced the signaling defect at physiological GM-CSF concentrations. Interestingly, at high GM-CSF concentrations similar to those observed in the index patient, signaling was partially rescued, thereby providing a molecular explanation for the slow progression of disease in these children. These results establish that GM-CSF signaling is critical for surfactant homeostasis in humans and demonstrate that mutations in CSF2RA cause familial PAP.

Pulmonary surfactant homeostasis is maintained by the balanced production of surfactant by alveolar epithelium and its clearance by pulmonary alveolar macrophages (1). GM-CSF is a critical regulator of surfactant homeostasis in mice (1). GM-CSF, via heterologous cell surface receptors comprised of α and β glycoprotein subunits (GM-CSF-Rα and GM-CSF-Rβ, respectively), activates STAT5 and other signaling pathways (2). N-linked glycosylation of the α subunit is required for GM-CSF binding and receptor signaling (3). In mice, deficiency of GM-CSF (4, 5) or its receptor (6) impairs multiple functions of alveolar macrophages (7), including surfactant catabolism (8), which results in progressive surfactant accumulation.

Pulmonary alveolar proteinosis (PAP) is a syndrome characterized by respiratory failure caused by pulmonary surfactant accumulation (9, 10), which can be grouped into distinct categories based on clinical, histopathologic, biochemical, and genetic data (1, 9–12). Primary PAP occurs when GM-CSF signaling is disrupted, for example in individuals with high levels of GM-CSF autoantibodies, which are
presumed to mediate pathogenesis by neutralizing GM-CSF and reducing surfactant catabolism by alveolar macrophages (10, 13). This form, referred to as autoimmune PAP, comprises 90% of cases (13). Primary PAP has also been associated with reduced detection of GM-CSF-Rβ on myeloid cells (14, 15), but definitive studies demonstrating heritable deficiency of either GM-CSF or its receptors as the cause of PAP in humans are lacking (16). Secondary PAP occurs as a consequence of an underlying disease presumed to impair surfactant clearance by reducing either the numbers or functions of alveolar macrophages (9). Hereditary disorders of surfactant production, for example, because of mutations in the genes encoding surfactant protein (SP)-B (17), SP-C (18), or ABCA3 (19), exhibit disordered surfactant homeostasis to varying degrees but are distinguished from PAP by their surfactant dysfunction, disruption of alveolar wall architecture, and clinical course (20). In this paper, we describe a family in which two children developed primary PAP in association with loss of GM-CSF responsiveness caused by mutations in the gene encoding GM-CSF-Rα.

RESULTS AND DISCUSSION

Clinical presentation and phenotype

The index patient presented at age six with a 2-yr history of progressive tachypnea and failure to thrive (height and weight were third percentile for age [Fig. S1 A, available at http://www.jem.org/cgi/content/full/jem.20080990/DC1]). Gestation, delivery, birth weight (3.66 kg, 50th percentile), and development were all normal, but weight gain slowed by 6–9 mo as did height by 2–3 yr. There was no history of cough, fever, chest pain, pneumonia or other pulmonary disease, environmental exposure, or drug use. Both parents were well-developed and healthy with no history of lung disease. Examination revealed moderate tachypnea, mild tachycardia, and inspiratory crackles but was otherwise unremarkable. Pulmonary function testing revealed severe restrictive impairment. Oxygen saturation was 88% while breathing room air and decreased while talking or walking a short distance. A diagnosis of PAP was suspected based on chest radiography (Fig. 1 A) and established by histopathologic examination of lung tissue (Fig. 1 B). A serum GM-CSF autoantibody test (21) was negative on two occasions. The patient was transferred to Cincinnati Children’s Hospital and underwent whole lung lavage therapy with marked symptomatic and radiographical improvement (Fig. 1 C). MCP-1 and M-CSF levels in the lavage were increased compared with healthy controls (supplemental text, available at http://www.jem.org/cgi/content/full/jem.20080990/DC1), which is similar to findings in autoimmune PAP patients and GM-CSF KO mice (22, 23).

An inherited defect in GM-CSF receptor function was suspected based on the constellation of histopathology showing well-preserved alveolar wall architecture (Fig. 1 B), the absence of GM-CSF autoantibodies (21), and an abnormal CD11b stimulation index test (24). Therefore, all immediate family members were evaluated. Because SP is in serum in PAP and levels reflect lung disease severity (25), serum SP-D levels were determined in the affected siblings and parents (Fig. 1 D). The mean ± SD serum SP-D concentration in 67 healthy controls was 63.5 ± 39 ng/ml (hatched region) and in 12 patients with autoimmune PAP was 174 ± 83 ng/ml (not indicated). The molecular basis for the observed increase in SP-D in these patients was examined by Southern blotting and by PCR of genomic DNA after digestion with restriction endonucleases (Table 1). Loss of heterozygosity was seen on chromosome 5q12-21 in all patients, consistent with inherited defects in GM-CSF-Rα and GM-CSF-Rβ (26).

Figure 1. Phenotypic characterization of patients with familial PAP. (A) Chest radiograph (top) and high-resolution computed tomogram of the chest (bottom) of the index patient at presentation. (B) Histopathologic appearance of the open lung biopsy from the index patient after staining with hematoxylin and eosin (H&E) or immunostaining for SP-A, mature SP-B, Pro–SP-C, SP-D, and ABCA3. Note the lymphocytosis in the low-power hematoxylin and eosin–stained section and the intact alveolar wall in the high-powered hematoxylin and eosin–stained section. Images were obtained at a total magnification of 50× (top) or 400× (all others). Bars: (top) 100 μm; (all others) 10 μm. (C) Chest radiograph of the index patient 4 mo after presentation immediately before (top) and 5 d after (bottom) whole lung lavage therapy. (D) Serum SP-D levels in affected siblings, parents, and a control. (E) High-resolution computed tomogram showing diffuse patchy ground glass opacities representing mild PAP in the affected sister.
was measured and found to be increased in the patient compared with her parents and controls (Fig. 1 D). Unexpectedly, the patient’s 8-yr-old sister, who was previously thought to be healthy, had increased serum SP-D. Subsequent clinical evaluation revealed that the sister had poor growth (height and weight, 10th percentile for age; Fig. S1B), a diffusion capacity for carbon dioxide of 57% that predicted, and mild patchy ground glass opacities throughout both lungs, which are consistent with a diagnosis of PAP (Fig. 1 E).

**Structure and function of the GM-CSF receptor**

Flow cytometry indicated that GM-CSF-Rα and GM-CSF-Rβ were present on blood leukocytes in all family members (Fig. 2 A). Western blotting revealed an abnormal electrophoretic pattern for GM-CSF-Rα in the patient and her sister with predominantly lower molecular mass forms compared with the control (Fig. 2 B). The father was heterozygous for normal and lower molecular mass forms. Unexpectedly, the mother exhibited only a normal pattern. To evaluate cell-mediated clearance of GM-CSF, blood mononuclear cells from the patient (Fig. 2 C) and her sister (Fig. S2, available at http://www.jem.org/cgi/content/full/jem.20080990/DC1) were incubated with GM-CSF, and both failed to clear GM-CSF in contrast to control and parental leukocytes. The concentration of GM-CSF in lung lavage was markedly increased in the patient compared with healthy controls, in whom it was undetectable (Fig. 2 D). The in vivo concentration of pulmonary GM-CSF in the epithelial lining fluid of the lungs was estimated to be 6.9 and 27 ng/ml (right and left lungs, respectively) using the urea dilution method (26).

**Figure 2. Structural and functional analysis of GM-CSF receptor defects.** (A) Flow cytometry demonstrating GM-CSF-Rα and GM-CSF-Rβ on the cell surface of peripheral blood leukocytes in all family members. (B) Western blots of PBMC lysates using antibodies to detect GM-CSF-Rα or actin as indicated. (C) GM-CSF clearance assay. Blood leukocytes from the patient were unable to remove exogenous GM-CSF added to culture medium at time zero in contrast to leukocytes from two controls that rapidly bound and cleared GM-CSF. Error bars show the means ± SE. (D and E) Measurement of GM-CSF concentration by ELISA. GM-CSF was readily detected in lavage from the patient but was not detected (ND) in lung lavage from nine healthy controls (D). GM-CSF was detected in the serum of affected siblings but not their parents or a control (E). (F) Blood leukocytes isolated from the indicated family members were incubated for 15 min in the absence (−) or presence (+) of 10 ng/ml GM-CSF, followed by Western blotting of cell lysates to detect total STAT5 (STAT5), phosphorylated STAT5 (pSTAT5), or actin. (G) Measurement of the GM-CSF-stimulated increase in CD11b levels on leukocytes in whole blood. The CD11b stimulation index (24) was calculated as the mean fluorescence of CD11b on leukocytes incubated with GM-CSF minus that of unstimulated cells divided by that of unstimulated cells and multiplied by 100. (H) Similar to F except that a higher GM-CSF concentration (1,000 ng/ml) was used and cell lysates were immunoprecipitated with anti-STAT5 antibody before Western blotting to detect total STAT5 (STAT5) or phosphorylated STAT5 (pSTAT5).
Figure 3. Genetic analysis of CSF2RA gene defects. (A) Nucleotide sequence of CSF2RA in genomic DNA from each family member, including nt 580–591 of the coding sequence (numbered relative to the initiation codon; from GenBank/EMBL/DDBJ under accession no. NM_006140.3). The index patient and her sister exhibited only a G→A point mutation at nt 586. The father was heterozygous for this substitution and the mother exhibited only the normal sequence. (B) FISH analysis to detect CSF2RA sequences in genomic DNA from the father and the patient. The probe (CTD-3047L21), which maps to the pseudoautosomal region (Xp22.33 and Yp11.32), hybridized to both X and Y chromosomes in the father (white arrows) and to one (white arrow), but not the other (yellow arrow), X chromosome in the patient. 8–10 metaphase cells and 25 interphase cells were evaluated for each individual. Similar FISH analyses are shown for each family member in the supplemental material (available at http://www.jem.org/cgi/content/full/jem.20080990/DC1). Images were obtained at a total magnification of 1000×. (C) CGH analysis for the patient in the region of Xp22.33. The relative fluorescence of fluorescently labeled DNA from the patient (open circles) compared with a same-sex reference DNA (filled circles) after hybridization to various BAC clones on the SignatureSelect V2 chip representing the Xp22.33 region is shown. Reduced hybridization to several BAC clones (clones B and C) is indicated by the lower fluorescence of the patient’s DNA compared with the reference DNA for these BAC clones. The telomeric breakpoint is mapped to between clones A and B and the centromeric breakpoint is mapped to between clones C and D. These data indicate an interstitial deletion of ~1.264 Mb at Xp22.33 encompassing ~1,610,183–2,873,864 bp. The dashed line represents a relative fluorescence of zero. (D) High-resolution SNP mapping of the Xp22 region for paternal and maternal X chromosomes. A schematic shows the locations of the point mutation (CSF2RA<sup>G174R</sup>) in the paternal X chromosome and the 1.6-Mb deletion at Xp22.33 in the maternal X chromosome and the genes encompassed. The base 2 ratio of normalized hybridization intensities for patient and reference samples (log R ratio) is shown. Similar SNP analyses for each family member are shown in the supplemental material (available at http://www.jem.org/cgi/content/full/jem.20080990/DC1). (E) Map showing a portion of the X chromosome summarizing the genetic analysis used to identify the small maternal X chromosomal deletion at Xp22.33 encompassing CSF2RA. The probe used for FISH analysis (hatched bar) is the same as clone B on the CGH microarray chip. The locations of selected CGH microarray probes in the region of the CSF2RA gene are shown. Those CGH probes showing
GM-CSF was also increased in CSF2RB-deficient mice (Fig. S3), confirming the relationship between GM-CSF receptor dysfunction and increased pulmonary GM-CSF levels. Serum GM-CSF was increased in both affected siblings and undetectable in both parents and control, indicating that the abnormality was systemic (Fig. 2 E). GM-CSF receptor function was impaired in both affected siblings, as shown by reduced GM-CSF-stimulated phosphorylation of STAT5 (Fig. 2 F) and failure of GM-CSF to increase CD11b levels on leukocytes from the patient compared with controls (Fig. 2 G).

The slower-than-expected rate of progression of PAP in both affected siblings led us to hypothesize that the GM-CSF-Rα abnormality may not completely abolish GM-CSF signaling. This was confirmed by incubation of primary blood mononuclear cells with a higher concentration of GM-CSF (1,000 ng/ml), which resulted in partial receptor signaling as shown by an increase in phosphorylated STAT5 (Fig. 2 H). Thus, although both GM-CSF-Rα and β chains were present on leukocytes in all family members, in the two affected siblings, the GM-CSF-Rα chain was structurally abnormal, which severely reduced but did not completely abolish GM-CSF binding and GM-CSF–dependent signaling and cellular functions.

Genetic analysis of the GM-CSF receptor

The nucleotide sequence of the CSF2RA gene in genomic DNA from both siblings revealed a single G→A point mutation in exon 7 that encoded a nonconservative amino acid change (glycine to arginine) at position 174, G174R (Fig. 3 A). This change alters one of the 11 potential sites of N-glycosylation, which is required for GM-CSF receptor function (3, 27). The G174R mutation present on the paternal X chromosome, explaining the unexpected CSF2RA G174R abnormality in one maternal X chromosome (Fig. 3 D) and in both siblings (Fig. S7). Importantly, this deletion encompassed the CSF2RA allele (Fig. 3 E), thereby explaining the unexpected CSF2RA DNA and mRNA sequence results.

These results demonstrate the inheritance of compound heterozygous abnormalities of CSF2RA in both siblings (Fig. 3 F). The G174R mutation present on the paternal X chromosome, localized to the extracellular cytokine binding domain of GM-CSF-Rα, alters a potential site of glycosylation and impairs GM-CSF binding and receptor signaling. However, it is unclear if the functional impairment is caused by the change in primary amino acid sequence or loss of glycosylation.

Reproduction of the CSF2RA<sup>G174R</sup> abnormality

To reproduce the GM-CSF receptor signaling defect identified in this family, plasmids expressing CSF2RB and either CSF2RA<sup>G174R</sup> or CSF2RA<sup>WT</sup> were cotransfected into 293 cells. Western blotting showed receptor proteins of the expected size from CSF2RA<sup>WT</sup> and smaller molecular forms from CSF2RA<sup>G174R</sup> (Fig. 4 A), which is similar to results for primary leukocytes (Fig. 2 B). Treatment of transduced cell lysates with Peptide: N-Glycosidase F, an amidase that cleaves away N-Linked glycosylation, confirmed the G174R mutation alters N-linked glycosylation (Fig. 4 A) by disrupting a functional N-linked glycosylation site and that glycosylation is responsible for the smaller molecular forms from this allele. GM-CSF-Rβ was expressed in CSF2RB-transfected cells but was undetectable in nontransfected cells (unpublished data). GM-CSF binding studies demonstrated that cells expressing CSF2RA<sup>WT</sup> and CSF2RB rapidly bound GM-CSF and removed it from the media, whereas cells expressing CSF2RA<sup>G174R</sup> and CSF2RB alone
did not clear GM-CSF (Fig. 4 B). Cotransfection of CSF2RB together with CSF2RA<sup>WT</sup>, but not with CSF2RA<sup>G174R</sup>, resulted in expression of functional GM-CSF receptors as demonstrated by GM-CSF-dependent phosphorylation of STAT5 only with the former combination (Fig. 4 C). Incubation of CSF2RA<sup>G174R</sup>/CSF2RB-transduced cells with higher concentrations of GM-CSF in the range of those observed in the index patient gave results similar to those obtained with primary leukocytes and clearly demonstrated a modest dose-dependent increase in STAT5 phosphorylation, which is consistent with a partial rescue of the signaling abnormality (Fig. 4 D).

**Clinical relevance of the CSF2RA<sup>G174R</sup> mutation**

Our findings demonstrate that familial primary PAP in humans is caused by compound heterozygous abnormalities of CSF2RA<sup>G174R</sup>, including loss of heterozygosity and a function-altering point mutation that severely reduces GM-CSF receptor signaling. These are the first results to demonstrate familial primary PAP in humans and that GM-CSF-Rα is critical for surfactant homeostasis. They extend previous findings that PAP is caused by the absence of GM-CSF or GM-CSF-Rβ in mice (4–6) or is specifically associated with the presence of neutralizing GM-CSF autoantibodies or a reduction in alveolar macrophage numbers or function in humans (9, 10). The allelic frequency of these mutations has not been determined but may be rare given that autoantibodies against GM-CSF account for most individuals with PAP.

Both affected siblings had impaired pulmonary function and failure to thrive, supporting the conclusion that prolonged respiratory insufficiency in children retards growth. Important to this conclusion is the observation that Xp22.33 deletion did not include the SOX (short stature homeobox) gene, the deletion of which impairs growth (28). Further, heterozygous loss of CSF2RA in the mother did not impair growth. Early results suggest that whole lung lavage therapy was followed by an increase in growth acceleration in the index patient.

This study highlights the usefulness of several novel biomarkers in the diagnostic evaluation of PAP. The combination of a negative GM-CSF autoantibody test (29) and an abnormal CD11b stimulation index (24) predicted the GM-CSF receptor dysfunction, which was confirmed by the lack of GM-CSF binding and defective GM-CSF receptor signaling. GM-CSF levels were increased in lung lavage from the patient and, in serum from the patient and her sister, compared with family members and healthy controls. Because primary PAP is presumed to be caused by a defect in GM-CSF regulation of alveolar macrophage surfactant catabolism (9, 10), an elevated serum GM-CSF level may be useful as a biomarker to distinguish primary PAP caused by GM-CSF autoantibodies (90% of all cases of PAP) (9, 10, 13, 30) from PAP caused by GM-CSF receptor dysfunction (this study; references 14–16) and PAP caused by GM-CSF deficiency (4, 5) (not yet reported in humans). Finally, an increased serum SP-D level predicted the occurrence of PAP in the sister, which was confirmed by subsequent clinical and radiographical evaluation.

The slow progression and differences in severity of the lung disease between affected siblings with the same CSF2RA mutations and GM-CSF-Rα dysfunction indicates that penetrance is important in familial PAP because of GM-CSF receptor defects. The increased levels of GM-CSF likely reflect impaired clearance but are also consistent with interruption of a
negative-feedback regulatory loop. Results strongly suggest that the high levels of pulmonary GM-CSF present in epithelial lining fluid result in a reduced but not abrogated receptor signaling and provide a molecular explanation the slow progression of PAP in both affected siblings. It is currently unclear if the level of increased pulmonary GM-CSF correlates with disease severity. They also suggest that aerosolized delivery of GM-CSF to the lungs may be therapeutic because this approach could theoretically achieve pulmonary GM-CSF levels several thousand–fold higher than endogenous levels measured in the patient and 20–fold higher than levels that appeared to completely restore GM-CSF receptor signaling in the transfection experiments. BM transplantation and CSF2RA gene transfer represent other potentially therapeutic approaches.

The identification of mild PAP in the older sister provides a unique and important opportunity to observe PAP at a very early stage of disease development. Finally, this case exemplifies the utility of a collaborative approach to the diagnosis and treatment of individuals with very rare diseases involving geographically disparate clinical centers and the benefit derived from integrating basic science, clinical medicine, and translational research.

MATERIALS AND METHODS

Participants. The institutional review board of the Cincinnati Children’s Hospital Medical Center approved the study, and all participants gave written informed consent or assent. Detailed case histories and laboratory data for the participants are included in the supplemental text (available at http://www.jem.org/cgi/content/full/jem.20080990/DC1).

GM-CSF receptor structure and function. GM-CSF receptors were evaluated by flow cytometry using antibodies recognizing GM-CSF-Rα (BD) or β (Santa Cruz Biotechnology, Inc.) and by Western blotting (7) with chain-specific antibodies (Santa Cruz Biotechnology, Inc.). The effects of glycosylation on GM-CSF-Rα were evaluated by incubating cell lysates with PNGase F (New England Biolabs, Inc.) (3). Actin was measured as a loading control using anti-actin antibodies (Santa Cruz Biotechnology, Inc.). To evaluate receptor-mediated clearance of GM-CSF, Ficol-purified blood leukocytes or 293 cells (American Type Culture Collection) were cultured of human GM-CSF Leukine for 15 min. After lysing RBC, leukocytes were evaluated by Western blotting (24) using anti–human STAT5 (Santa Cruz Biotechnology, Inc.) and anti- phospho-STAT5 (Millipore) antibodies. The authors have no conflicting financial interests.

ELISA. SP-D, GM-CSF, MCP-1 (monocyte chemotactic protein-1), M-CSF, and urea were measured using commercial ELISA kits (BioVendor, R&D Systems, BD, R&D Systems, and BioAssay Systems, respectively) (24). The concentration of cytokines in epithelial lining fluid of the lungs was estimated by multiplying the concentration in lavage fluid by the ratio of urea concentrations in serum and lavage fluid (26).

Genetic analysis. Genomic DNA or total mRNA was purified from blood leukocytes with commercially available kits (QIAGEN). The PCR was used to generate products spanning exons and flanking intronic sequences (genomic DNA) or the coding sequences and 5’ and 3’ flanking untranslated sequences (mRNA) of the CSF2RA and CSF2RB genes, which were then analyzed by direct sequencing. The resulting sequences were compared with published sequences (from GenBank/EMBL/DDBJ under accession nos. NM_006140.3 and AC070885, respectively). The relative copy number of CSF2RA and CSF2RB genes among family members and a healthy control was measured by amplification of genomic DNA using PCR with gene-specific oligonucleotides followed by analysis of products by electrophoresis on 2% agarose gels. Experimental methods for measuring gene copy number, evaluating the genes encoding SP-B, SP-C, and ABCA3 (19, 31) for function-altering mutations, cytogenetic analysis, CGH analysis, FISH, and high-resolution SNP array analysis (32) are described in the supplemental Materials and methods (available at http://www.jem.org/cgi/content/full/jem.20080990/DC1).

Reproduction of the GM-CSF receptor defect. The methods used to clone and express mRNA transcripts encoding GM-CSF-Rα and β chains in cultured cells and evaluate protein glycosylation are described in the supplemental Materials and methods.

Statistical analysis. Numeric data are presented as means ± SE. Statistical comparisons were made with Student’s t test for two-group comparisons and with one-way analysis of variance with posthoc analysis according to the Holm-Sidak method for multiple-group comparisons. P-values <0.05 were considered to indicate statistical significance.

Online supplemental material. Fig. S1 shows normalized growth curves for the index patient and her sister. Fig. S2 shows GM-CSF binding and clearance by blood leukocytes from affected and unaffected family members and control. Fig. S3 shows increased pulmonary GM-CSF levels in GM-CSF receptor β chain (Csf2rb) KO mice. Fig. S4 shows evaluation of CSF2RA and CSF2FB gene copy number by PCR amplification. Fig. S5 shows karyotypes of high-resolution GTG-banded chromosomes. Fig. S6 shows evaluation of heterozygosity at the CSF2RA locus by FISH. Fig. S7 shows high-resolution SNP analysis. The supplemental text contains additional details regarding the case histories and clinical laboratory and other data cited throughout the text. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20080990/DC1.

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REFERENCES


PARTICIPANTS

Patient. The index case was a 6-yr-old girl who was admitted to the hospital in July, 2007 when a chest radiograph revealed bilateral diffuse alveolar infiltrates. She was the product of a normal gestation and delivery and had reached developmental milestones as expected. At birth, she weighed 3.64 kg and was 19.5 cm in length (both at the 50th percentile); however, by 6–9 mo weight gain slowed and by 2–3 yr height acceleration slowed (Fig. S1 A). On admission, both her weight (13.6 kg) and height (103.5 cm) were below the third percentile for her age. She was noted by her mother to avoid swimming underwater at 3–4 yr of age, refusing to hold her breath. By 4–5 yr of age, she had developed dyspnea of insidious onset, first with exercise and then at rest, and finally was noted to “pant” in her sleep. There was no history of cough, fever, chest pain, lymphadenopathy, pneumonia or serious infection, pulmonary or other disease, environmental exposures, or drug use. There was no family history of lung disease in children or unexplained deaths at an early age. On examination, the respiratory rate was 60 per minute, blood pressure was 88/30 mm Hg, and pulse was 106. Pulmonary auscultation revealed mild crackles during deep inspiration, but the remainder of the exam was otherwise unremarkable.

The white blood cell count was 5.9 × 10⁹/ml with normal cytometric indices, the RBC count was 4.9 million/mm³, the hemoglobin was 13.9 g/dcliter, and the hematocrit was 41.1%. The levels of serum IgG, IgA, and IgM were mildly elevated (995, 150, and 181 g/dcliter, respectively) and IgE was 6.2 g/dcliter. Pulmonary function testing revealed a pattern of severe restriction with a forced vital capacity (FVC) that was 44% of that predicted, forced expiratory volume in one second (FEV1) 44% that of predicted, and a FEV1/FVC ratio 116% of that predicted. Oxygen saturation was 88% while breathing room air and dropped when talking or walking a short distance. A chest radiograph showed diffuse patchy alveolar infiltrates throughout both lung fields (Fig. 1 A, top). Bronchoscopic examination of the airways was unremarkable. Cytological examination of the bronchoalveolar lavage fluid demonstrated the presence of lipoproteinaceous material. A serum GM-CSF antibody test was negative on two occasions. The urine did not contain detectable amino acids. No sequence deviations were detected in the coding exons and intron–exon boundaries of the genes encoding SPs B and C and ABCA3 (1, 2).

The patient was transferred to Cincinnati Children’s Hospital in October, 2007 and underwent whole lung lavage therapy, which resulted in marked symptomatic improvement (Fig. 1 C). 4 mo later, the oxygen saturation was 98% on room air, and pulmonary function testing revealed an FVC that was 69% of that predicted, an FEV1 of 61% that predicted, and an FEV1/FVC ratio of 116% that predicted. The weight had increased to 16 kg and the height to 107.5 cm. Exercise tolerance had markedly improved and the patient resumed more normal daily activities.

Sister. The sister of the index patient was 8 yr old in February, 2008 when first evaluated after a negative GM-CSF autoantibody test and an abnormal CD11b stimulation index test suggested that an inherited GM-CSF receptor defect may be present in the index case (3). She was the product of a normal gestation and delivery and had reached developmental milestones as expected. At birth, her weight and length were 3.6 kg and 20 cm (both at the 50th percentile) and at the time of evaluation were 22.7 kg and 122 cm, respectively (both at the 10th percentile for age; Fig. S1B). She had been considered by her family to have been healthy all her life with no history of lung disease or major illnesses. On examination, the respiratory rate was 32 per minute, the pulse was 82 per minute, and blood pressure was 104/62 mm Hg and was otherwise unremarkable.

Pulmonary function testing demonstrated an FVC that was 88% of that predicted, FEV1 that was 90% of that predicted, a FEV1/FVC ratio 116% of that predicted. Oxygen saturation was 97% while breathing room air. An HRCT of the chest revealed mild patchy ground glass opacities throughout both lungs, which is consistent with the diagnosis of PAP (Fig. 1 E). The severe reduction in DLCO and failure to thrive suggest that the lung abnormalities are more extensive than indicated by the chest HRCT; i.e., scattered ground glass opacifications involving geographically distributed secondary pulmonary lobules. Rather, these results suggest that a thickened alveolar surfactant layer may be present throughout the alveolar surface but is not well visualized by the radiographical technique used.

Father. The father of the index patient was 49 yr old at the time of evaluation in February, 2008. His height was 185.4 cm (85th percentile for his age). He was a lifelong nonsmoker with no history of pulmonary or other major medical illness.

Mother. The mother of the index patient was 48 yr old at the time of evaluation in February, 2008. Her height was 165 cm
(65th percentile for her age). She was a lifelong nonsmoker with no history of pulmonary or other major medical illness.

**Patients with autoimmune PAP.** 12 individuals with autoimmune PAP are reported as disease controls, all of whom had the typical clinical, physiological, and radiographical features of the disease at the time of evaluation. The detailed case histories and diagnostic criteria of each have been reported previously (3). The group included three pediatric and nine adult patients. The mean ± SE GM-CSF autoantibody titer was 324.3 ± 66 µg/ml and was similar in adult and pediatric cases (365 ± 74 and 202 ± 141 µg/ml, respectively; P = 0.308).

**Healthy controls.** Volunteers were enrolled in the study as healthy controls. This control group included 67 individuals, 61 of whom were previously reported as part of another study (3). The mean ± SE age was 30 ± 6 yr. All were disease-free healthy individuals without a history of major illness and all were symptom-free at the time of enrollment in the study. None were current smokers.

**MATERIALS AND METHODS**

**Lung histopathology and immunostaining**

A formalin-fixed paraffin-embedded open lung biopsy tissue sample was received for evaluation. 5-µm-thick sections were cut on a rotary microtome and loaded onto polylysine-coated slides. Hematoxylin and eosin-stained sections were prepared using routine methods. Immunohistochemistry was performed as described previously (4, 5) using rabbit polyclonal antisera raised against SP-A, SP-B, pro–SP-C, and SP-D, a mouse monoclonal antibody generated to ABCA3 as primary antibodies and biotinylated secondary antibodies (Vector Laboratories), and an avidin–biotin–horseradish peroxidase detection system (ABC reagent; Vector Laboratories). The enzymatic reaction product was enhanced using Ni-DAB and Tris cobalt to give a black precipitate, and the sections were counterstained with Nuclear Fast Red.

**GM–CSF clearance assay**

To evaluate receptor-mediated binding of GM–CSF, heparinized blood was obtained by phlebotomy and mononuclear cells were isolated on Ficoll gradients, followed by RBC lysis as previously described (3). Primary mononuclear cells or transfected 293 cells (see Construction of vectors and transfection) were seeded into culture dishes at 1 × 106 cells per dish in DMEM containing 10% bovine calf serum. Human GM–CSF (Leukine; Bayer) was added at a concentration of 1 ng/ml. At subsequent times, GM-CSF was measured in aliquots of the culture medium by ELISA (R&D Systems). The results were expressed as a percentage of initial GM-CSF concentration.

**Nucleotide sequence analysis**

The nucleotide sequence of CSF2RA and CSF2RB transcripts and genomic DNA were determined by PCR-based methods in the Genetic Variation and Gene Discovery Core Facility at the Cincinnati Children’s Hospital Medical Center. Genomic DNA was also evaluated for the presence of known function-altering mutations in the coding sequences and intron–exon boundaries of the genes encoding SP-B, SP-C, and ABCA3 as previously described (1, 2).

**Measurement of gene copy number by PCR amplification**

The relative copy number of CSF2RA and CSF2RB genes among family members and controls was determined using genomic DNA using PCR with the following gene-specific primers: 5′-AGGAG AAGAGGGAGATCCG-3′, 5′-CACGTG-GCCTCAGTTCACAG-3′ (CSF2RA); 5′-ACAG AGCCAGGCAATGTTGTT-3′, and 5′-CGACAAACTCCTGG-CGAGG-3′ (CSF2RB). Amplification conditions were as follows: 94°C for 5 min; 94°C for 30 s, 58°C for 30 s, and 72°C for 45 s (30 cycles); and 72°C for 7 min. PCR-generated products were then evaluated by electrophoresis on 2% agarose gels.

**Cytogenetics**

Chromosome analysis was performed on cells isolated from peripheral blood according to standard cytogenetic techniques.

**Comparative genomic hybridization**

Comparative genomic hybridization analysis was performed with genomic DNA isolated from each family member using the SignatureSelect V2 microarray chips (Signature Genomic Laboratories, LLC) and a fluorescence dye reversal assay (6). Microarray chips were processed according to the manufacturer’s protocol. Genomic DNA were evaluated with a same-sex reference DNA (patient) or an opposite sex reference DNA (mother and sister) and data analysis was performed using the Genoglyphix software (Signature Genomic Laboratories, LLC).

**Fluorescence in situ hybridization**

FISH analysis was performed under routine clinical laboratory protocols with the CTD-3047L21 hybridization probe, which identifies the pseudoautosomal region of X and Y chromosomes, Xp22.33 and Yp11.32, respectively. Slides were analyzed with a fluorescent microscope (Carl Zeiss, Inc.) with CytoVision v3.7 software (Applied Imaging), 8–10 metaphases and 25 interphase cells were evaluated in each FISH study.

**High-resolution single-nucleotide polymorphism analysis**

Blood was obtained from the patient, her sister, and parents, and genomic DNA was isolated using a commercial kit (QIAGEN) according to the manufacturer’s instructions. Samples of genomic DNA (750 ng) from each subject were prepared for microarray analysis using the Infinium Assay (Illumina Inc.) according to the manufacturer’s protocol. Microarray analysis was performed using the HumanCNV370-duo DNA Analysis BeadChip platform (Illumina Inc.). This chip contains ~370,404 markers, including SNP and copy number variation content. Data were analyzed using the BeadStudio v3.1 analysis software (Illumina Inc.).

**Construction of vectors and transfection**

PBMC RNA from the index patient and a healthy control was purified and converted to cDNA as described in the Ma-
terials and methods section Genetic analysis. The nucleotide sequence for the entire protein coding sequence of CSF2RA was generated from cDNA transcripts by PCR amplification using gene-specific primers and subcloned into pSC-A (Agilent Technologies). After confirming the nucleotide sequence, a restriction fragment containing the entire coding region of CSF2RA from the control (CSF2RAWT) or the index patient (CSF2RAG174R) were inserted individually into a plasmid (MIEG3) permitting expression in mammalian cells (7). The cDNA encoding CSF2RB was obtained (IART98-E2; Geneservice) and inserted into another plasmid (MSCV2.1) permitting expression in mammalian cells (8). Expression plasmids carrying CSF2RWT and either CSF2RAG174R or CSF2RAT were cotransfected into human embryonic kidney epithelial cells (HEK 293 cells; American Type Culture Collection) using Lipofectamine2000 (Invitrogen) according to the manufacturer’s instructions. After 48 h, cells were evaluated for (1) expression of GM-CSF receptor proteins by Western blotting (2), GM-CSF binding, and clearance by culturing with exogenous GM-CSF (see GM-CSF clearance assay) or (3) GM-CSF receptor signaling by evaluation of GM-CSF–dependent phosphorylation of STAT5 (see STAT5).

Western blotting

GM-CSF-Rα was evaluated in blood leukocytes or transfected cells as follows. Leukocytes from 1 ml of blood (evaluated within 1 h of phlebotomy and subjected to RBC lysis) or 1 × 10⁶ transfected 293 cells were lysed in 200 µl RIPA buffer (0.05 M Tris-HCl, pH 8, 0.15 M NaCl [Tris-buffered saline (TBS)], 1% vol/vol nonidet P-40, 0.5% wt/vol sodium deoxycholate, 0.1% wt/vol SDS, 0.004% wt/vol sodium azide) containing 2% vol/vol protease inhibitor cocktail, 1% vol/vol phenyl-methyl-sulfonyl-fluoride (PMSF), and 1% vol/vol sodium orthovanadate (Santa Cruz Biotechnology, Inc.) as directed by the manufacturer. Samples were kept on ice for 30 min and then centrifuged (9,000 g at 4°C for 15 min) to remove insoluble debris. Samples were mixed with equal volume of sample buffer and boiled. The lysate was then fractionated on SDS-PAGE gels (4–12% Tris Glycine gel; Invitrogen) under reducing conditions and proteins were transferred to PVDF membranes by electroblotting. Membranes were incubated in blotting solution (TBS, 5% [wt/vol] dry milk, and 0.1% [vol/vol] Tween 20) at 4°C, overnight to block nonspecific binding. Mouse antihuman GM-CSF-Rα antibody (diluted 1:500; Santa Cruz Biotechnology, Inc.) was then added and incubations were continued at room temperature for 2 h. After washing in TBS and 0.1% (vol/vol) Tween 20, membranes were incubated at room temperature for 1 h in blotting solution containing donkey anti–rabbit IgG. After washing in blotting solution, membranes were incubated with ECL-Plus (GE Healthcare) to visualize immunostained proteins as directed by the manufacturer. This procedure was used for measuring actin in the same samples with anti-Actin antibody (diluted 1:1,000; Santa Cruz Biotechnology, Inc.).

STAT5.

GM-CSF receptor signaling was evaluated in blood leukocytes or transfected cells as described for GM-CSF-Rα with the following modifications. 1 ml of blood (evaluated within 1 h of phlebotomy) or 1 × 10⁶ transfected 293 cells (in DMEM containing 10% bovine calf serum) were incubated in the absence or presence of human GM-CSF (Leukine; 1, 10, 100, and 1,000 ng/ml) for 15 min at 37°C. After RBC lysis (whole blood) and washing with PBS (both leukocytes and 293 cells), cell lysates were prepared and evaluated as in GM-CSF-Rα, except that for transfected 293 cell lysates and cells exposed to high GM-CSF concentrations, immunoprecipitation was performed at 4°C overnight using ProteinA/G Agarose (Santa Cruz Biotechnology, Inc.) and anti–STAT5 antibody or control rabbit IgG (Santa Cruz Biotechnology, Inc.). Anti–STAT5 antibody (diluted 1:500; Santa Cruz Biotechnology, Inc.) or anti–phospho-STAT5 (diluted 1:500; Millipore) were used for primary antibodies for Western blotting.

Evaluation of CSF2RA glycosylation

Expression plasmids carrying either CSF2RAG174R or CSF2RAT were transfected into 1 × 10⁶ HEK 293 cells as described in Construction of vectors and transfection. After 48 h, cells were lysed in 200 µl RIPA buffer containing protease inhibitor cocktail, PMSF, sodium orthovanadate (Santa Cruz Biotechnology, Inc.). Samples were kept on ice for 30 min and then centrifuged at 9,000 g at 4°C for 15 min to remove insoluble debris. Nine µl of lysate was treated with PNGase F (New England Biolabs, Inc.) then added and incubated at 37°C for 30 min according to the manufacturer’s protocol and then evaluated by Western blotting as in GM-CSF-Rα.

Measurement of cytokine concentration in lung epithelial lining fluid recovered by bronchoalveolar lavage

The volume of epithelial lining fluid (ELF) recovered in lung lavage fluid was estimated using the urea dilution method described by Rennard et al. (9). In brief, the concentration of urea in the lung lavage fluid and serum were measured by ELISA. The volume of ELF (in milliliters) was calculated as the volume of lavage fluid multiplied by the concentration of urea in lavage fluid in milligrams per milliliter divided by the concentration of urea in plasma in milligrams per milliliters. The concentration of cytokines in ELF was estimated by multiplying their concentration in lavage fluid (measured by ELISA) by the ratio of the lavage fluid in milliliters to ELF volume in milliliters.

REFERENCES


