Hermansky-Pudlak syndrome type 2 manifests with fibrosing lung disease early in childhood

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Abstract

Background: Hermansky-Pudlak syndrome (HPS), a hereditary multisystem disorder with oculocutaneous albinism, may be caused by mutations in one of at least 10 separate genes. The HPS-2 subtype is distinguished by the presence of neutropenia and knowledge of its pulmonary phenotype in children is scarce.

Methods: Six children with genetically proven HPS-2 presented to the chILD-EU register between 2009 and 2017; the data were collected systematically and imaging studies were scored blinded.

Results: Pulmonary symptoms including dyspnea, coughing, need for oxygen, and clubbing started 3.3 years before the diagnosis was made at the mean age of 8.83 years (range 2-15). All children had recurrent pulmonary infections, 3 had a spontaneous pneumothorax, and 4 developed scoliosis. The frequency of pulmonary complaints increased over time. The leading radiographic pattern was ground-glass opacities with a rapid increase in reticular pattern and traction bronchiectasis between initial and follow-up Computer tomography (CT) in all subjects. Honeycombing and cysts were newly detectable in 3 patients. Half of the patients received a lung biopsy for diagnosis; histological patterns were cellular non-specific interstitial pneumonia, usual interstitial pneumonia-like, and desquamative interstitial pneumonia.

Conclusions: HPS-2 is characterized by a rapidly fibrosing lung disease during early childhood. Effective treatments are required.

Keywords: Hermansky-Pudlak syndrome type 2, Childhood, Pulmonary fibrosis, Tachydyspnea, Pulmonary phenotype

Background

Hermansky Pudlak syndrome is a rare hereditary multisystem disorder first described in 1959. More than 50% of all worldwide cases are identified in individuals from Puerto Rico where HPS has an estimated frequency of 1:1800 [1, 2]. Clinically the syndrome is characterized by oculocutaneous albinism, a bleeding diathesis due to platelet storage deficiency [1, 2], and other manifestations which may include neutropenia, a granulomatous colitis, or pulmonary fibrosis.

Genotypic analysis over the last decade allowed differentiating at least 10 separate forms of HPS, due to mutations in different genes [3]. All entities have in common defects in intracellular protein trafficking and the biogenesis of lysosome-related organelles like melanosomes or platelet dense granules [2].

Pulmonary fibrosis has not been described in HPS-3 and HPS-5 through HPS-10, which are all very rare. HPS-1 is the most common subtype and characteristically develops a severe and progressive pulmonary fibrosis in almost all cases. Usually middle-aged adults and rarely late adolescents are affected by fibrosis; however, so far, children are not described [2-4]. HPS-4 has been documented in less than 10 patients, few of which had pulmonary fibrosis [5, 6].
The HPS-2 subtype is also very rare, with less than 40 cases reported worldwide [4, 7–10]. HPS-2 is caused by mutations in the AP3B1 gene, inherited in an autosomal recessive way and distinguished from the other forms of HPS by the presence of neutropenia that can lead to severe respiratory infections and that is responsive to granulocyte colony-stimulating factor [1]. Among the few patients described, development of an interstitial lung disease (ILD) has been mentioned in 30 to 50% [1, 7]; details on the pulmonary phenotype have been described in four cases [4]. Potential mechanisms causing pulmonary disease in HPS-2 are poorly understood. It has been suggested that altered AP3B1 gene product within alveolar epithelial type II cells leads to defective intracellular processing of surfactant proteins B and C (SP-B, SP-C), endoplasmic reticulum-stress, apoptosis, and a fibrotic lung phenotype [1].

The aim of this study was to describe the pulmonary phenotype of HPS-2 in children and to further investigate the presence and the possible role of cellular stress and apoptosis in patient-derived material.

Methods

Patients, diagnosis, and follow up

Patients were recruited from the chILD-EU register and biobank and the kids’ lung register collecting diffuse parenchymal lung diseases [11, 12]. Among the children included between 2009 and 2017, seven children were diagnosed with HPS. A 0.4-year old infant with HPS-1 referred for the assessment of potential pulmonary involvement had no pulmonary symptoms and was excluded from this study. All the other cases were HPS-2.

The diagnosis of HPS-2 was based on typical clinical symptoms and proven by genetic analysis (Table 1). Mutational analysis was performed by Sanger sequencing. Routine clinical evaluation in different European centers was performed; data were collected retrospectively and prospectively following the inclusion into the study.

Lung function testing was done according to standards set previously in children old enough to perform spirometry [13]. CT images of the chest were evaluated for the presence of parenchymal abnormalities (like mosaic attenuation, ground glass opacity, consolidation, linear opacity, septal thickening, reticular opacity, nodular opacity, honeycombing, emphysema, cysts, bleb or bulla) and airway abnormalities (tree-in-bud, bronchiectasis, bronchial wall thickening) on a lobar basis, counting lingula as the separate lobe [14]. Also the presence of pneumothorax, pleural thickening, pleural effusion and enlarged hilar or mediastinal lymph node were evaluated. The image analysis was performed blinded by a pediatric radiologist with expertise in chest imaging.

Bronchoscopy and bronchoalveolar lavage (BAL)

Flexible Bronchoscopy including BAL (mostly of the middle lobe) were performed if clinically indicated using 3 times 1 ml warmed normal saline per kilogram body weight. BAL was examined cytologically and microbiologically.

Lung biopsies and histological investigations

Lung biopsies available were peer-reviewed independently and blinded by a pathologist specialized in pulmonary pathology. Lung tissue of patient 3 was analyzed by Western blotting under reducing and denaturing conditions using sodium dodecyl sulfate–polyacrylamide gel electrophoresis followed by electroblotting and immunostaining for pro-SP-C (Merk Millipore, Darmstadt, Germany), ATF6, β-actin (abcam, Cambridge, UK), and cleaved caspase-3 (Cell Signaling, Gaithersburg, USA). Blotted membranes were developed with the ECL Plus chemiluminescent detection system (Amersham Biosciences, Amersham, UK). Immunohistochemistry was performed on lung tissue fixed in 4% formaldehyde on serial sections with the AP Fast Red kit (Zytochem Systems, Berlin, Germany) after antigen retrieval by microwaving in 10 mM sodium citrate buffer, pH 6.0. Hemaluna was used as counter-stain. Slides from patient 2 were also available for immunostaining for pro-SP-C and cleaved caspase-3, as described above. As controls, lung sections from 3 different organ donor lungs were used.

Ethics, consent and permissions, consent to publish, declarations and statement

Informed consent to report individual patient data was obtained by all patients old enough to consent, and their parents or guardians. The study was approved by the ethics committee of the Ludwig-Maximilian University of Munich (EK 111-13).

All supporting data have been entered in the additional material (Additional file 1).

Result

Clinical course and mutations detected

The six patients included had a mean age at diagnosis of 8.83 years (Additional file 1: Detailed description of the individual cases. Subjects 1 to 6). On average, lung symptoms started 3.3 years before diagnosis (Table 2, Additional file 1: Table S1). Most frequent signs and symptoms reported at follow-up were dyspnea, coughing, need for oxygen, tachypnea, and clubbing. At last follow-up on average 3.17 years (range 2-6) after diagnosis, pulmonary symptoms were noted in all patients. Overall, the frequency of all pulmonary complaints increased over time. Non-pulmonary signs and symptoms are listed in Additional file 1: Table S1.

Patients 1 and 5 were siblings and had the same homozygous frameshift mutation. All other children had
### Table 1 Baseline demographics and genetics

|                      | Patient 1          | Patient 2          | Patient 3          | Patient 4          | Patient 5          | Patient 6          | Distribution   |
|----------------------|--------------------|--------------------|--------------------|--------------------|--------------------|--------------------|----------------|
| Gender               | female             | female             | male               | female             | female             | female             | female         |
| Consanguinity by history | yes                | no                 | yes                | no                 | yes, sibling of patient 1 | no; genealogy demonstrated common ancestors, i.e. very distant relation | female |
| Alleles\textsuperscript{a} | homozygous         | homozygous         | homozygous         | compound heterozygous | homozygous         | homozygous         | 5: 1, homozygous: compound heterozygous |
| \(AP3B1\) mutation 1 | c.3222-3223delTG (\(\sim\)) | g.151312_159483del8172bp (\(\sim\)) | c.2546 T > G (\(\sim\)) | c.177delA (\(\sim\)) | c.3222-3223delTG (\(\sim\)) | c.2944delC (\(\sim\)) | 2 known, 4 previously unknown variants |
| \(AP3B1\) mutation 2 | c.3222-3223delTG (\(\sim\)) | g.151312_159483del8172bp (\(\sim\)) | c.2546 T > G (\(\sim\)) | c.1839_1842delTAGA (\(\sim\)) | c.3222-3223delTG (\(\sim\)) | c.2944delC (\(\sim\)) | unknown, likely pathogenic variant (frame shift; early termination in exon 26) |
| Previously described; predicted pathogenic effect of mutations | known [10]; likely pathogenic variant (frame shift) | known [28]; likely pathogenic variant (exon skipping) | unknown; likely pathogenic variant (stop-mutation; early termination in exon 22 instead of 27) | unknown; likely pathogenic variant (frameshift mutations and subsequent early terminations in exons 2 and 17) | known [10]; likely pathogenic variant (frame shift) | unknown, likely pathogenic variant (frame shift; early termination in exon 26) | unknown |

\textsuperscript{a}All parents tested were heterozygous for the respective variants; not tested in patient 3. \textsuperscript{b}GenBank References (NM_003664.4) and \textsuperscript{c}(NG_007268.1)
different mutations in AP3B1, two of the variations were described before. All other mutations are likely disease-causing mutations, as they are predicted to result in truncated AP3B1 either through the introduction of a premature stop signal, by omission of a regular stop codon or by skipping of an important structural domain for adaptor protein 3 (AP-3 complex) formation. (Table 1, Fig. 1). Although a history of consanguinity could not be obtained in all cases, only one patient was compound heterozygous for the mutations, supporting the autosomal recessive nature of the disease.
pattern of inheritance (Table 1). Of interest, 5 of 6 patients were girls. Siblings with the same mutation had different clinical courses. However, the small size of this cohort precludes definite conclusions about genotype-phenotype associations in patients with HPS-2.

Complications related to the respiratory tract
In 2 patients scoliosis was noticed at diagnosis, together with psychomotor retardation. Scoliosis developed during follow-up in 2 additional patients. In 3 patients, spontaneous pneumothorax occurred during follow-up. Pleurodesis was necessary for 2 patients (Table 2). In patients old enough to perform lung function testing, a slightly impaired diffusion capacity for carbon monoxide was noted. Initially, spirometry was in the mild abnormal range with combined restrictive-obstructive ventilation disorder, with changing course over time (Table 3, case descriptions in Additional file 1 and Additional file 2).

Chest imaging
CT scans were performed at the time of diagnosis and at follow-up in subjects 1, 3 and 6 (Fig. 2, Additional file 3: Figure S1, Additional file 4: Figure S2, Additional file 5: Figure S4, Additional file 6: Figure S5, Additional file 7: Figure S6). For subject 4, CT was performed only at the time of diagnosis and for subject 2 only at follow-up. Age at first CT scan was 6.4 years (range 2.3-12) and 12.9 years (range 5.3-15.4) at follow-up. Leading radiographic pattern at diagnosis was ground-glass opacity, which had a patchy distribution, occurring in almost each lobe in all patients. In patient 1 also reticular opacities and traction bronchiectasis were detected at diagnosis. Patient 3 had bronchial wall thickening in the initial CT scan. There was a rapid increase in reticular pattern and traction bronchiectasis between initial and follow-up CT in all subjects with two examinations. Honeycombing and cysts were newly detectable in 3 patients after an observation time of 5.2 years. No nodular opacity, consolidation, tree-in-bud, emphysema, or air trapping were detected (Additional file 1: Table S2, Fig. 2, Additional file 3: Figure S1, Additional file 4: Figure S2, Additional file 5: Figure S4, Additional file 6: Figure S5, Additional file 7: Figure S6). There was no predominance of one side; upper lobes showed the most frequent affection.

BAL and histology
Four of 6 patients had a BAL at diagnosis. One child had very mild eosinophilia and neutrophilia. Patient 6 had severe neutrophilia, despite peripheral blood neutropenia, suggesting compartmentalized capacity to mobilize neutrophils into the alveolar space (Additional file 1: Table S3). No pathogenic bacteria were recovered; patients were not on antibiotic treatment at the time of diagnosis.

Table 3 Lung function measurements, lung biopsy results, treatments, and overall outcome

|                         | Patient 1 | Patient 2 | Patient 3 | Patient 4 | Patient 5 | Patient 6 | Mean (n) or distribution |
|-------------------------|-----------|-----------|-----------|-----------|-----------|-----------|-------------------------|
| Age [y] at first / last lung function | 12 / 17 | nd (psychomotor retardation) | 9 / 19 | nd (too young) | nd (psychomotor retardation) | 8/15 | 9.67 / 17 (3) |
| FEV1 [% predicted] first / last | 74 / 71 | 82 / 66 | 93 / 59 | 62/67 | 73 / 70 (3) |
| FVC [% predicted] first / last | 65 / 70 | 74 / 72 | 69 (15 y)/ 71 (19 y) | 67.75 (4) |
| TLC [% predicted] | nd / 80 | nd | cNSIP, UIP-like, DIP UIP-like | nd | cNSIP, DIP, lymphoid follicular hyperplasia | 3 biopsies |
| Lung biopsy | nd | cNSIP, UIP-like, DIP | UIP-like | nd | nd | cNSIP, DIP, lymphoid follicular hyperplasia |
| Chronic antibiotic treatment | Co-trimoxazole | Azithromycin | no | Azithromycin | Azithromycin | no | 4/6 |
| G-CSF s.c. | yes | yes | no | yes | yes | yes | 5/6 |
| Pirfenidone treatment | yes (started age 13 for 30 months) | no | yes (age 15 for 3 months) | no | no | no | 2/6 |
| Age last follow-up [y] | 17 | 14 | 19 | 3.7 | 15 | 12.45 (range 3.7-19) |
| Overall outcome | sick-better | sick-same | sick-same | sick-same | died | sick-same | 4 sick-same, 1 sick-better, 1 died |

Abbreviations: y year(s), FEV1 forced expiratory volume of first second, FVC forced vital capacity, TLC total lung capacity, DLCO diffusing capacity of the lung for carbon monoxide, cNSIP cellular non-specific interstitial pneumonitis, nd not done, UIP-like usual interstitial pneumonia-like features, DIP desquamative interstitial pneumonitis, G-CSF granulocyte-colony stimulating factor
Three of the 6 patients received a lung biopsy for diagnosis. Due to bleeding diathesis and low thrombocyte values, thoracoscopic biopsies were performed instead of transbronchial biopsies. In patient 2, the histological pattern of cellular non-specific interstitial pneumonitis (cNSIP), usual interstitial pneumonia-like features (UIP-like), and of desquamative interstitial pneumonitis (DIP) were noted and correlated to the pattern identified on chest CT (Additional file 4: Figure S2). Patient 3 had an UIP-like pattern of patchy dense fibrosis with subpleural cystic areas, pleural fibrosis, and pleural blebs. Histology showed hyperplastic and vacuolated type II pneumocytes and ceroid containing macrophages as typical features of HPS (see Fig. 3a-d). Patient 6 underwent lung biopsy at the age of 6 years. Comparable to patient 2 the histological pattern of cNSIP with areas of DIP were noted. Moreover, lymphofollicular hyperplasia with few lymphoid follicles was detectable. Typical vacuolated type II pneumocytes cells were found ubiquitously but only few ceroid containing macrophages (Additional file 8: Figure S7).

Treatment
Because of neutropenia and resulting immune deficiency, chronic antibiotic treatment in 4 and Granulocyte-Colony Stimulating Factor, (G-CSF) treatment in 5 patients were initiated. The latter resulted in a lower normal range of neutrophil peripheral blood counts. Pirfenidone was used in 2 patients but was stopped after 3, respectively 24 months, due to gastrointestinal side effects and lack of sufficient clinical improvement (Table 3).

Alveolar epithelial cellular stress and apoptosis
In patient 3 with frozen tissue available obtained from 2 different parts of the lung, additional mechanistic investigations were performed and compared to healthy tissue obtained from unused lung of organ donors. The myofibroblast marker α-smooth muscle actin (SMA) was increased in one specimen supporting increased fibrosis (Fig. 4). Following whole lung tissue analysis, we further analyzed type II pneumocyte apoptosis, an important event in the pathogenesis of interstitial lung diseases. Serial sections revealed several type II pneumocytes to be positive for the apoptosis marker, cleaved caspase 3, indicating their apoptosis (Fig. 3e, Additional file 4: Figure S2H). Two types of cellular stress, the endoplasmic reticulum (ER) stress and autophagy can be differentiated in type II pneumocytes of HPS type 1 patient lungs [15, 16]. We thus analyzed the ER stress markers, GRP78 and p50 form of ATF6, which were markedly increased in one and moderately in the other specimen of the lungs of the HPS-2 patient (Fig. 4). Similarly, the autophagosomal marker microtubule-associated protein
1 light chain-\(3\beta\), LC3B in its lipidated form (LC3BII) was elevated compared to two of the control lungs, one also gave an increased signal, along with a concomitant increase in the autophagy substrate protein, p62. This proved a defective autophagy pathway in the lung of this HPS-2 patient (Fig. 4a).

**Discussion**

Children suffering from HPS-2 may present with a severe and progressive chronic pulmonary phenotype. Severe lung fibrosis may develop until early adulthood; associated complications including pneumothorax, recurrent infections, and the development of scoliosis were key features identified. Together with few forms of ATP-binding cassette sub-family A member 3 (ABCA3) \[17–19\], SFTPC \[20, 21\], and MARS deficiency \[22\], this condition operates under the few clearly and rapidly fibrosing diffuse parenchymal lung diseases in childhood.

Clinically HPS-2 in children is diagnosed by the combination of albinism, bleeding diathesis, and neutropenia. The evolution of symptoms in 4 children started with epistaxis or bleeding diathesis as initial symptoms during the first 2 years of life. However, respiratory symptoms were present in all our children but obviously were rated as too non-specific or developed too insidious to contribute to the diagnosis at age 5 years. At that time, half of our patients already had clubbing, dyspnea, and oxygen demand. Seventeen of the 22 cases under 18 years of age in the largest series on HPS-2 so far had respiratory symptoms, although not further specified \[7\]. Tachypnea and wet coughing developed somewhat later and suggest secondary chronic bronchitis or suppurrative lung disease. The latter may be due to additional immune deficiency from neutropenia in HPS-2 subjects, which may aggravate early respiratory affections in these children. Three of our 6 patients developed pneumothoraces, adding substantially to pulmonary morbidity. Subpleural lung fibrosis, in particular cysts or honeycombing, predisposes to such a complication, which is very unusual in children at this young age. Among 4 children with HPS-2, Gochuico et al. described one child with 6 recurrent pneumothoraces \[4\].

The natural history of HPS-2 lung disease differs from that of HPS-1 pulmonary fibrosis, which usually affects
middle-aged adults and not children [23]. Based on published reports, patients generally first manifest symptoms of HPS pulmonary fibrosis in middle age, however, anecdotal experience includes rare patients with HPS-1 beginning to develop ILD in late adolescence [24]. Carmona-Rivera described a 16-year-old boy with no pulmonary symptoms in HPS-1 [25]. Characteristic pulmonary CT findings known in adults with HPS-1 are increased reticular opacities, thickened interlobular septa, and ground-glass infiltrates in addition to fibrotic changes, including traction bronchiectasis, subpleural cysts and honeycombing. These imaging findings evolve over time, starting in early adulthood, as in HPS-1 patients < 20 years usually no CT changes are noted, in those between 20 and 29 years minimal changes were identified, with increasing symptoms in patients 30 years and older [26, 27]. In HPS-1, high-resolution CT abnormalities inversely correlated with percentage of forced vital capacity and were useful in defining the progression of interstitial disease [27]. In our patients with HPS-2, patchy ground-glass opacity predominated at diagnosis during early childhood and a reticular pattern evolved rapidly over time. At follow-up most of the children developed the CT findings characteristic for HPS fibrosis in adults. Thus, compared to patients with HPS-1, in HPS-2 not only pulmonary symptoms as described above but also CT abnormalities were detected very early. Of interest and in contrast to our observations, two children with HPS-2 have been described in literature before with bilateral ground-glass opacity, thickening of interlobular septa, and interstitial reticulations (4 and 8 vs. 14 years of age) and 3 more children suffering from HPS-2 had changes in HRCT all with improvement over time [4, 9].

In all children, the diagnosis of HPS-2 was verified genetically (Table 1). A pair of siblings had the same mutation (patient 1 and 5), however their clinical course was different. This was also due to the fact that the pulmonary...
phenotype of the second child was much more and earlier in focus after the other died. A frameshift mutation in the C-terminal region of \( AP3B1 \) resulted in loss of the stop codon, prolonging translation into the 3'UTR region. Although an alternative in-frame stop codon is available further downstream, increased distance to the original stop codon may predispose transcripts to nonstop-mediated decay mechanisms [28]. Alternatively, the translated protein product might be subject to proteolytic breakdown due to misfolding, defective assembly, or intrinsic conformational instability [29]. Patient 2 displayed a larger genomic deletion which includes parts of introns 14, 15, and exon 15. This specific region has shown to be essential for correct assembly of the AP3-complex [30]. In all other individuals, single point mutations (patient 3) or frameshift mutations caused by base pair deletions about 10-120 base pairs upstream (patient 4, 6) lead to premature stop codons, translation termination, and possibly activation of the nonsense-mediated decay pathway (Fig. 1).

The pulmonary fibrosis in patients with HPS may be preceded by a macrophage-mediated alveolar inflammation, as BAL fluid contains increased numbers of constitutively activated macrophages [26]. Only one of the subjects with BAL had increased macrophage counts (Additional file 1: Table S3). All our patients had progressively fibrosing lung disease already during childhood. Our histological data prove that pulmonary fibrosis affects children with HPS-2. This is in contrast to HPS-1 and 4, where the development of pulmonary fibrosis starts in middle-age adults and children are only rarely affected [24]. Overall rapidly progressive fibrosing lung disease in childhood is extremely rare. The development of fibrosis might be facilitated by the patients’ neutropenia and natural killer - and T-cell dysfunction and the resulting susceptibility to severe recurrent chest infections. Such an observation is consistent with the HPS-2 animal model where environmental lung injury by silica or bleomycin aggravates fibrosis [26]. Therefore, preventive measures as vaccination and aggressive antibiotic treatment are warranted. Taking our limited observation length into account, the clinical course was not stable in most children. Despite intense symptomatic treatments after diagnosis, we saw deterioration and development of complications in several patients. Four patients remained unchanged, one patient improved, one patient (subject 5), however, died from respiratory insufficiency.

The histology of HPS-2 interstitial lung disease is not widely explored, as the diagnosis may now be done genetically. Patient 2 had a combination of NSIP and UIP-like pattern with dense fibrosis in peribronchiolar and subpleural distribution, as well as a DIP-like areas with intraalveolar aggregates of alveolar macrophages, very similar to other descriptions published [4]. Lung biopsy of patient 3 was dominated by a patchy dense fibrosis with UIP-like pattern comparable to patient 2. Lung biopsy of patient 6 did not show areas of dense fibrosis but demonstrated a cellular NSIP pattern. Additionally, there was a mild lymphoid hyperplasia with few lymphoid follicles with germinal centers that could probably be interpreted as post-infectious changes.

Apoptosis of type II pneumocytes, in addition to ER stress and defective autophagy, was observed in a HPS-2 patient lung. This finding is in line with the previously reported observations of cellular stress and apoptosis of type II pneumocytes in several interstitial lung diseases. More studies are needed to determine if defective autophagy or ER stress underlie type II pneumocyte apoptosis and are subsequently responsible for fibrotic remodeling in the HPS-2 patient lung.

**Conclusion**

We highlight that patients with HPS-2 in contrast to the other forms of HPS must be considered a severely and rapidly fibrosing lung disease already during early childhood which requires effective antifibrotic treatment. The latter is not yet available in childhood. At diagnosis, half of the subjects had clinical signs of chronic hypoxemia. Histology remains of importance in those extreme rare diseases, which should be made based on the clinical phenotype and confirmed by targeted genetics. During the course, which may be complicated by recurrent pneumothoraces and scoliosis, a comprehensive multidisciplinary team approach is needed. Currently early symptomatic pulmonary care including consequent antibiotic treatment of pulmonary infections and vaccinations. Nutritional management and early orthopedic treatment of scoliosis may help to improve the outcome of children with HPS-2. No specific antifibrotic treatment is available in children.

**Additional files**

**Additional file 1:** Detailed description of the individual cases. Subjects 1 to 6. Table S1. Non-pulmonary signs and symptoms. Table S2. Blinded scoring of CT-scans (number of CTs available at diagnosis (n = 5)/ at follow-up (n = 5); CTs at follow-up and diagnosis available in sub 1-3 and 6, only at follow-up in sub 2 and only at diagnosis in sub. 4). Scans were analysed for the presence and absence of features listed in the six sections of the lungs. Scoring was not validated. Table S3. Results of bronchoalveolar lavage (BAL) at diagnosis. (DOC 107 kb)

**Additional file 2:** Figure S3. BODYPLETHYSMOGRAPHY OF SUBJECT 3. Bodyplethysmography at the last follow up (19 years of age). Visible is the restrictive pattern with mild obstruction. (TIFF 807 kb)

**Additional file 3:** Figure S1. CT scan and spirometry course of subject 1. (A) CT scan at 12 and 15 years of age. Initially patchy GGO (black arrowheads) and mild reticulations were seen in all lobes with apical predominance, at follow-up also reticular pattern increased and traction bronchiolitis and mild honeycombing appeared (black arrow). (B) Spirometry long-term course of PVC, which shows variabilities of 15% during the last 4 years. Pirfenidone treatment led to improvement first, but...
Figure S2. Additional file 4: Figure S2. CT scan and histological pattern of subject 2. (A) CT scan at the age of 13 8/12 years. CT images show the thorax deformity due to scoliosis with displacement of the central bronchial structures and traction of the segmental bronchi. On the left side, cystic parenchymal destruction in the apex (black arrow), mild bronchial wall thickening in the upper lobe (encircled), reticular opacities predominantly in the upper lobe and lingula. Increased lung density especially in the lower lobe is due to breathing artefacts and areas of ground-glass opacity. On the right side, there are few subpleural cysts in the apex and reticular opacities in the upper and middle lobe. (B-G) Histological pattern with patchy areas of dense interstitial fibrosis (red) in peribronchial and subpleural distribution (B; HE, x10). Dense fibrosis with small areas of honeycombing with bronchiolar metaplasia (C; HE, x50). Fibroblastic foci as features of UIP pattern (D; HE, x 200) and areas with NSIP pattern consisting of diffuse mild interstitial fibrosis and a mild chronic inflammatory infiltrate in between (E; HE, x100). Few small interstitial aggregates of ceroid macrophages (F; HE, x200) and vacuolization of hyperplastic type II pneumocytes (G; HE, x400). (H) Increased in the apoposis marker cleaved caspase-3 in alveolar epithelial cells type II (AECII) in HPS2 patient lungs. Representative images of immunohistochemistry for cleaved caspase-3 and proSP-C (AECII marker) performed on serial paraffin sections of HPS2 and donor lungs. Shown here are high magnification images, 40x objective for both HPS2 and donor lung sections. Arrows indicate same AECII stained for both cleaved caspase-3 and SP-C implying dying AECII in HPS2 patient lungs while almost no cleaved caspase-3 is detected in SP-C positive AECII of donor lungs. Scale bar, as indicated in the images. (TIF 4239 kb)

Additional file 5: Figure S4. CT scan of subject 4. CT Scan with patchy distribution of ground glass opacity throughout all lobes. (JPEG 8908 kb)

Additional file 6: Figure S5. CT scan of subject 5. CT at 2 (left column) and 5 (right column) years of age. Initial scan was performed in prone position, follow up in supine position. Initial scan demonstrates linear opacities (black arrows) with mild traction bronchiectasis. After a course of 3 years CT scan deteriorates with severe parenchymal abnormalities showing diffuse ground glass opacity, reticular opacity, mild honeycombing, and cysts (black arrowheads). Moreover enlarged mediastinal and hilar lymph nodes have been seen (better seen on the mediastinal window, images not provided here). (TIF 7675 kb)

Additional file 7: Figure S6. CT scan of subject 6. CT images at 6 (first column), 9 (second column), 11 (third column) and 15 (fourth column) years of age. At the initial scan subject presented with patchy GGO in all lobes and additional interlobular septal thickening (encircled in white). At first follow up, a decrease of GGO in the left upper and lower lobes can be seen (black arrows in initial scan, first column). Over the next years, the distribution and extent of GGO showed no relevant changes. The extents of interlobular septal thickening decreased at the first follow up, followed by increase of reticular markings (encircled in black) in the next scans. Also bronchial wall thickening was present on the last follow up scans, here best seen in the right lower lobe in the third column (dashed circle). (TIF 4655 kb)

Additional file 8: Figure S7. Histological pattern of subject 6. A (HE, x50). Normal lung architecture with partly collapsed alveolar spaces on the right containing aggregates of alveolar macrophages (black arrows). Hypercellular interstitium with diffuse mild lymphocytic infiltrate in alveolar septa and few lymphoid follicles. B (HE, x100). Prominent lymphoid follicle with central germinal center, interstitial lymphocytes and intravascular macrophages. C (HE, x200). Aggregates of intravascular macrophages with pink cytoplasm (black arrows), interstitial lymphocytes with small blue nuclei (black triangles), scattered type II pneumocytes with clear vacuolated cytoplasm (white arrows) typically found in HPS. (TIF 5371 kb)

Abbreviations
ABC3A3: ATP-binding cassette sub-family A member 3; AP3: Adaptor protein 3; AP3B1: Adapter protein 3 complex subunit beta-1; ATF6: Activating transcription factor 6; BAL: Bronchoalveolar lavage; CNASP: Cellular non-specific interstitial pneumonitis; CT: Computer tomography; DIP: Desquamative interstitial pneumonitis; ER: Endoplasmic reticulum; G-CSF: Granulocyte-Colony Stimulating Factor; GRP78: Glucose-regulated protein 78; HPS: Hermansky-Pudlak syndrome; IILD: Intestinal lung disease; LC3B1: Light chain-3; LC3B in its lipidated form; MARS: Methionyl-tRNA Synthetase; mM: Millimolar; p50: Protein 50; p62: Protein 62; SMA: Smooth muscle actin; SP: Surfactant protein; UIP: Usual interstitial pneumonia

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Availability of data and materials
All supporting data have been entered in the additional material (Additional File 1). None of the authors has any competing interests in the manuscript.

Authors’ contributions
MH collected the data and drafted the manuscript. LN, JGO, STL, LHS, FB contributed and discussed cases, PM did biochemical and histological investigations, AG reviewed those data and the manuscript, LAS reviewed the genetic data, SR read all pathology, JLZ evaluated all imaging data, performed data interpretation, and reviewed the manuscript. MG designed the project, reviewed all cases, performed data interpretation, and reviewed the manuscript. All authors approved the final version of the manuscript.

Ethics approval and consent to participate
The study was approved by the ethics committee of the Ludwig-Maximilians University of Munich (EK 111-13).

Consent for publication
Informed consent to report individual patient data was obtained by all patients old enough to consent, and their parents or guardians.

Competing interests
The authors declare that they have no competing interests.

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