Familial idiopathic pulmonary fibrosis in association with bone marrow hypoplasia and hepatic nodular regenerative hyperplasia: a new “trimorphic” syndrome

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ABSTRACT
This is the first report of familial idiopathic pulmonary fibrosis associated with hepatic nodular regenerative hyperplasia and bone marrow hypoplasia. Four members of one family presented with this triad of organ dysfunction. The response to immunosuppressive treatment was poor and all four members succumbed to the disease processes. The current literature is reviewed and mechanisms that could have been involved in the development of this new syndrome are proposed.

We report the first case of familial idiopathic pulmonary fibrosis (FIPF) associated with hepatic nodular regenerative hyperplasia (NRH) and bone marrow hypoplasia. We have named this “trimorphic syndrome”—three (inherited) morbidities, pulmonary, hepatic and cytopenia.

CASE HISTORY
Sibling 1
A 34-year-old man presented with a 3-week history of cough and dyspnoea. He had no medical history, had never smoked and worked as a builder/carpenter. He was clubbed, had harsh breath sounds throughout both lung fields and hepatosplenomegaly. The chest radiograph showed ill-defined nodules throughout both lung fields. Pulmonary function tests (PFT) and high-resolution CT (HRCT) scan of his chest were consistent with pulmonary fibrosis (table 1).

The serum angiotensin-converting enzyme (ACE) level was normal. He was thrombocytopenic but bone marrow examination was normal (table 2). Liver enzymes were mildly raised (table 2), while ultrasonography demonstrated diffusely abnormal liver architecture and portal hypertension. A full liver and autoimmune screen (autoantibodies, immunoglobulins, viral hepatitis B and C serology, α1-antitrypsin, caeruloplasmin, celiac serology) was negative. Serum ferritin was mildly elevated (507 μg/l, normal range 15–300), but haemochromatosis gene (C282Y/H63D) mutation analysis was negative.

He deteriorated clinically and was commenced on high-dose prednisolone. Repeat bone marrow examination showed hypocellularity (table 2). He underwent a transjugular liver biopsy but deteriorated during the procedure and died (10 months after presentation). Post-mortem examination (tables 1 and 2) confirmed pulmonary fibrosis and NRH (fig 1).

Father
Five months later the 74-year-old father, a retired bricklayer, presented with a 4-month history of dyspnoea. Examination revealed inspiratory crepitations. PFTs were consistent with pulmonary fibrosis, while the HRCT scan showed fibrosis and pleural thickening (table 1). He was thrombocytopenic but liver enzymes were normal (table 2). Serum autoantibodies (nuclear and ANCA) were negative, with minimal elevation of serum ACE (59 IU/l, normal range 9–50).

He failed to attend appointments and died suddenly 18 months later. Post-mortem examination confirmed IPF (table 1), portal hypertension, NRH (table 2) and mild myocardial fibrosis.

Sibling 2
Four years after sibling one presented, his 46-year-old brother presented with a 12-month history of cough and dyspnoea. He had never smoked and worked as a bricklayer. Examination revealed bilateral inspiratory crepitations. PFTs and HRCT scan were consistent with pulmonary fibrosis (table 1).

The serum angiotensin-converting enzyme (ACE) level was normal. He was thrombocytopenic but bone marrow examination was normal (table 2). Liver enzymes were mildly raised (table 2), while ultrasonography demonstrated diffusely abnormal liver architecture and portal hypertension. A full liver and autoimmune screen (autoantibodies, immunoglobulins, viral hepatitis B and C serology, α1-antitrypsin, caeruloplasmin, celiac serology) was negative. Serum ferritin was mildly elevated (507 μg/l, normal range 15–300), but haemochromatosis gene (C282Y/H63D) mutation analysis was negative.

He deteriorated clinically and was commenced on high-dose prednisolone. Repeat bone marrow examination showed hypocellularity (table 2). He underwent a transjugular liver biopsy but deteriorated during the procedure and died (10 months after presentation). Post-mortem examination (tables 1 and 2) confirmed pulmonary fibrosis and NRH (fig 1).

Table 1 Pulmonary features at presentation by person

<table>
<thead>
<tr>
<th></th>
<th>PFTs</th>
<th>TcO (% predicted)</th>
<th>HRCT chest</th>
<th>Lung histology</th>
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<tr>
<td>Sibling 1</td>
<td>Restrictive pattern</td>
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<td>Father</td>
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<td>Fibrosis and pleural thickening</td>
<td>IPF*</td>
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<td>Widespread fibrosis and honeycombing</td>
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<td>Sibling 3</td>
<td>Normal limits</td>
<td>78</td>
<td>Patchy fibrotic change consistent with IPF</td>
<td>IPF*</td>
</tr>
</tbody>
</table>

*At post-mortem examination. †At surgical lung biopsy.
HRCT, high-resolution computed tomography; IPF, idiopathic pulmonary fibrosis; PFT, pulmonary function test; TcO, transfer factor for carbon monoxide.
confirmed by surgical lung biopsy (table 1). He was thrombocytopenic with a hypocellular bone marrow (fig 2), with areas of increased reticulin suggestive of early secondary fibrosis. Liver enzymes were mildly elevated (table 2). Serum ACE, rheumatoid factor, serum autoantibodies and liver screen were negative.

Despite treatment with prednisolone and azathioprine, he died 6 months after presentation. Post-mortem examination revealed IPF (table 1), hepatic NRH (table 2), pleural plaques and mild cardiac fibrosis.

Sibling 3
An asymptomatic brother aged 57 years was referred alongside sibling 2. He had never smoked and worked as a carpenter/joiner. Examination revealed scattered bi-basal inspiratory crepitations. PFTs were normal but HRCT of his chest was consistent with IPF (table 1). He was thrombocytopenic with low to normal marrow cellularity but normal liver function tests (table 2). Rheumatoid factor and autoantibodies were negative, while the serum ACE level was mildly elevated (84 IU/l, normal range 9–50).

Despite treatment with prednisolone and azathioprine, his lung function deteriorated. He developed raised liver enzymes despite cessation of azathioprine, with ultrasonography showing a shrunken liver and portal hypertension (table 2). A liver screen was normal but the ferritin level was raised (942 μg/l, normal range 15–300); he was negative for haemochromatosis gene mutations (C282Y/H65D). Repeat bone marrow examination showed worsening hypocellularity with areas suggestive of early fibrosis (table 2). He died 4 years after presentation; post-mortem examination showed IPF (fig 3) and NRH (table 2).

DISCUSSION
This is the first report of FIPF associated with hepatic NRH and bone marrow hypoplasia—a combination of fibrotic (lung) and non-fibrotic (liver and bone marrow) processes.

FIPF is defined as at least two members of a biological family (parent, child, sibling) with clinical features of IPF confirmed

<table>
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<th>Table 2</th>
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<td>Platelets $\times 10^9$/l</td>
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<td>Sibling 1</td>
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<tr>
<td>Father</td>
<td>98</td>
</tr>
<tr>
<td>Sibling 2</td>
<td>73</td>
</tr>
<tr>
<td>Sibling 3</td>
<td>87</td>
</tr>
</tbody>
</table>

*At post-mortem examination. †On ultrasound scan at presentation. ‡On ultrasound scan 3.5 years after presentation. 3.5 years after presentation.

**Figure 1** Liver biopsy showing a nodule of larger hepatocytes surrounded by compressed hepatocytes (reticulin stain, ×2 objective lens).

**Figure 2** Marrow trephine biopsy showing excess fat spaces with reduced haemopoietic precursors indicative of a hypoplastic marrow (H&E stain, ×20 objective lens).
by interaction between a gene (or genes) and an environmental trigger. All our cases had worked in the building trade and two as carpenters, with exposure to wood dust associated with IFP.

Two had also handled asbestos which can cause pulmonary fibrosis similar to IFP. Neither of these occupational exposures, however, provides a causal link to the hepatic and haematological abnormalities.

In sporadic IPF it is believed that micro-insults to alveolar epithelial cell walls are followed by aberrant wound healing, with fibroblasts depositing extracellular matrix in the distal air spaces. A number of cytokines, growth factors and other cell mediators are involved in this process, one or more of which must have systemic actions since bone marrow-derived fibrocytes are recruited to the lungs in IPF. Such systemic actions could explain the more widespread fibrotic process seen in family members with cardiac and marrow fibrosis and be responsible for the hepatic and haematological abnormalities seen in this family.

Hepatic NRH is characterised by hepatocellular nodules in the absence of fibrous septa between these nodules. Its main complication is portal hypertension. Occurring in 5.6% of individuals over 80 years, it is commonly associated with vasculitic conditions, neoplastic and haematological illnesses and drugs including azathioprine. Two of the cases received azathioprine but one already had abnormal liver enzymes when it was started.

Tissue ischaemia is believed to play an important role in its pathogenesis. In celiac disease IgA anticardiolipin antibodies are thought to trigger thromboses in portal vein radicles draining the small intestine, leading to liver damage and NRH. Such pathogenic factors might “spill over” into the lungs when the hepatic filtration system fails; interestingly, local activation of the coagulation cascade has been demonstrated in lung models of IPF. However, siblings 1 and 3 had normal anticardiolipin levels and neither had the common thrombophilia risk factors (gene mutations for factor V Leiden and prothrombin and functional deficiencies of protein C, protein S and antithrombin III). We are also unable to explain the haematological findings seen in our patients on the basis of such a hypothesis.

Bone marrow hypoplasia is a reduction in the number of haemopoietic progenitor cells in the marrow. It can be congenital, such as in Fanconi’s anaemia, or be acquired...
secondary to radiation, viral infections,\textsuperscript{9} drugs\textsuperscript{10} and immune-mediated conditions such as systemic lupus erythematosus.\textsuperscript{11} It can be seen in myelofibrosis where marrow is replaced by fibrous tissue. Features typical of myelofibrosis were not, however, seen in these cases; in contrast, mild patchy fibrosis was seen postdating the hypocellularity.

Patients with NRH often present with thrombocytopenia, thought to be secondary to platelet consumption within hepatic thrombi or to undiagnosed hypersplenism from portal hypertension. However, they would be expected to develop a hyperplastic marrow rather than the hypoplastic marrow seen in our cases.

Sibling 2 was thrombocytopenic when he underwent a minor surgical procedure 18 months before presentation (platelets 98 $\times 10^3$/l). This might suggest that a bone marrow-derived factor or the absence of a factor normally secreted by the marrow is responsible for both the hepatic and lung pathologies, perhaps causing liver microvascular congestion and nodule formation (ie, NRH) and lung fibrosis through local activation of coagulation cascades. However, such mechanisms remain speculative.

In summary, we have described a new syndrome of familial lung fibrosis, bone marrow hypoplasia and hepatic NRH. Although no unifying pathological process has been identified, we have discussed putative factors. As there are no known genotypic markers for FIPF, screening of the remaining family members is limited to phenotype surveillance. We have offered the remaining sibling and future offspring of the index generation 5-yearly chest radiography, full blood count and ultrasound scan of the liver. Treatment options, however, remain limited as all affected members have demonstrated a poor response to medical treatment.

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**Competing interests:** None.

**Patient consent:** Obtained from relatives of the patients.

**REFERENCES**

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