Horizons in Medicine: Update in diagnosis and management of interstitial lung disease

<table>
<thead>
<tr>
<th>Journal</th>
<th>Clinical Medicine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID</td>
<td>CM-2016-0265.R1</td>
</tr>
<tr>
<td>Manuscript Type</td>
<td>Review</td>
</tr>
<tr>
<td>Keywords</td>
<td>Interstitial lung disease, Idiopathic pulmonary fibrosis, cryoscopic lung biopsy</td>
</tr>
</tbody>
</table>
ABSTRACT

The field of Interstitial Lung Disease (ILD) has undergone significant evolution in recent years, with an increasing incidence and more complex, ever expanding disease classification. In their most severe forms these diseases lead to progressive loss of lung function, respiratory failure and eventually death. Despite notable advances, progress has been challenged by a poor understanding of pathological mechanisms and patient heterogeneity, including, variable progression. The diagnostic pathway is thus being continually refined, with the introduction of tools such as transbronchial cryo lung biopsy and a move towards genetically aided, precision medicine. In this chapter, we will focus on how to approach a patient with ILD and the diagnostic process.

INTRODUCTION

Interstitial lung disease (ILD) is an umbrella term for over 200 different diseases which display considerable variation in terms of clinical course, treatment and prognosis. Broadly speaking they can be sub-divided into those with an identifiable cause and those without, the latter being referred to as idiopathic interstitial pneumonias (IIPs). Clinical assessment aims to identify a possible cause; screening for features of systemic disease (e.g. connective tissue disease, CTD) or environmental triggers. Relevant exposures include pneumotoxic drugs, radiation therapy, occupational exposures (e.g. asbestosis) or implicated allergens (hypersensitivity pneumonitis, HP).

Distinguishing the various forms of pulmonary fibrosis is critical for determining correct management and for predicting prognosis, however, this is often obfuscated by the fact the lung has a limited repertoire in response to injury and consequently, a finite number of disease patterns. In essence, all ILD is characterised by variable degrees of inflammation and fibrosis, not only between diseases, but also among individuals with the same disease (Figure a). In inflammation dominant disease the histology is that of organising pneumonia, or non-specific interstitial pneumonitis (NSIP), whilst in fibrosis dominant disease, one would expect to see usual interstitial pneumonitis (UIP), characterised by fibroblastic foci and only mild to moderate inflammation. These histological patterns are associated with specific radiological features, the recognition of which may abrogate the need for a formal biopsy and tissue diagnosis.
IDIOPATHIC PULMONARY FIBROSIS

IPF is the most common of the IIPs, with an incidence of approximately 6,000 cases per year in the UK, affecting mainly older males. Median survival is worse than many cancers at just three years and the disease accounts for 5,300 deaths each year in the UK (1). IPF is a growing problem, with an annual increase in incidence of 11% between 1991 and 2003, a rise that is only partly explained by an ageing population (2).

As already alluded to, distinguishing IPF from other ILD, including other idiopathic forms, is important for multiple reasons. IPF does not respond to immunosuppressive therapy, in fact, immunomodulation may worsen outcomes (3). By contrast there is evidence, particularly in systemic sclerosis associated ILD (4), of benefit from cyclophosphamide and multiple case reports suggest a potential role for rituximab (5) as salvage therapy in CTD-ILD. In addition, there are now two drugs, pirfenidone and nintedanib, approved by NICE for IPF, however, at an annual cost of around £26,000 per patient and the potential for significant side effects, accurate disease identification is essential. Finally, IPF has a worse prognosis than other ILD, hence a definitive diagnosis allows for timely involvement of palliative care physicians and consideration of lung transplantation.

PATHOGENESIS OF IPF

The pathogenesis of IPF is complex and poorly understood, but involves aberrant wound healing in the context of repetitive alveolar injury. This results in abnormal fibroblast proliferation, differentiation and activation, which in turn drives expansion of the extracellular matrix with loss of normal lung architecture, whilst inflammation plays a less dominant role. This is illustrated schematically in Figure a.

GENETICS IN DIAGNOSIS AND MANAGEMENT OF IPF

Although the initiating events in IPF are poorly understood, the disease is likely to be the result of environment exposures in genetically susceptible individuals. Certainly, it is estimated that approximately 20% of IIPs have a genetic component and familial cases (referred to as Familial Interstitial Pneumonias, FIP) were first described in the 1950s. The majority of these FIPs are autosomal dominant with incomplete penetrance, but some may arise de novo. The most commonly affected genes are those involved in surfactant processing and telomere biology. At present, routine genetic testing is not recommended, however, ILD patients with at least one affected first degree
relative, should be offered the opportunity to enrol in the UK wide 100,000 Genomes Project, whereby they undergo whole genome sequencing.

Genetics also have a proven role in sporadic IPF. Polymorphisms in the promoter for the gene encoding the salivary mucin, 5b (MUC5B) and for the Toll-interacting protein (TOLLIP) are both associated with an increased risk of developing IPF, although both result in a relatively mild phenotype. These genetic variants provide the first possible targets for precision medicine in IPF, with a post hoc analysis of the PANTHER data establishing a variable response to N-acetyl cysteine depending on the individual’s TOLLIP genotype (6).

DIAGNOSTIC WORK UP IN INTERSTITIAL LUNG DISEASES

Initial investigations include blood tests to detect the presence of autoantibodies (Table 1), precipitating immunoglobulins against organic antigens and serum ACE. These tests alone rarely confirm the diagnosis and there is potential for both false positive (particularly autoantibodies in older patients) and false negative (failure to identify an antigen and IgG does not exclude HP) results, however, they can be useful in helping direct further diagnostics.

Pulmonary function tests (PFTs) are key in appraising these patients and whilst they rarely refine the specific diagnosis in individuals with proven ILD, they inform on disease severity at baseline and response to treatment over follow up.

RADIOLOGICAL WORK UP

Chest Radiographs
A Chest x-ray (CXR) is often the first radiological investigation in ILD patients and whilst it is rarely sufficient to make a confident diagnosis, X-ray can play a role in establishing disease chronicity and progression.

High Resolution Computed Tomography
High-resolution computed tomography (HRCT) of the thorax has revolutionised the diagnosis and classification of ILD and in many cases removes the need for invasive diagnostic procedures, however, the quality of the images is dependent on the scanning protocol employed (Table 2).

The ATS/ERS 2011 consensus statement (7) provides criteria for a definite UIP pattern on HRCT (Figure c), with the presence of these conferring a sensitivity of approximately 40%, but a specificity of 95% for histological UIP. The main discriminating feature for UIP is the presence of
honeycombing, however, typical CT appearances are only present in two thirds of patients and it is in the remaining third of cases, that biopsy may have a role.

BRONCHOSCOPY IN THE ASSESSMENT OF ILD

Bronchoalveolar Lavage, Endobronchial Ultrasound and Transbronchial Biopsy

There is much variation in practice surrounding the use of bronch-alveolar lavage (BAL), particularly between European centres and North American colleagues, who rarely utilise it. Clearly, there is value in excluding infection, which may be a differential diagnosis, however, BAL alone is rarely diagnostic, with perhaps one of the difficulties being a lack of consistency in terms of how samples are taken and processed. Under optimal circumstances BAL reflects cellular traffic in the alveolar space and the cell differential may provide supplemental information to help refine, rather make a diagnosis. In particular, an excess of lymphocytes should call into question a presumptive IPF diagnosis, with Ohshimo et al. describing a BAL lymphocytosis of >30% in 6 of 74 patients with definite UIP features on HRCT. In all six cases, further investigations led to a final diagnosis of chronic hypersensitivity pneumonitis (8).

Transbronchial biopsy (TBB) with standard forceps is a minimally invasive technique, but does not always provide adequate lung tissue to establish a final diagnosis. The biopsies are small, subject to crush artefact and may not be representative in spatially heterogeneous disease (9,10). It may, however, be helpful in the diagnosis of sarcoidosis and organising pneumonia.

Transbronchial Cryo Lung Biopsy

Transbronchial cryo lung biopsy (TBCLB) was first described in 2008 (11). It has since been shown to be a safe, minimally invasive and effective diagnostic tool for the histological diagnosis of ILD, with a diagnostic yield of up to 74–80% (11–15). The advantage of TBCLB over TBB lies in the larger specimen size, with a mean size range of 9mm$^2$ to 64mm$^2$ (11–20). In addition, the technique avoids crush or bleeding artefact, which can distort the tissue architecture (Figure d). The published data on TBCLB shows a safety profile that is comparable to TBB, with bleeding post biopsy in around 10% of cases, all of which was controlled bronchoscopically. The mean rate of pneumothorax requiring chest drain insertion is around 4%, although there is a wide variation between centres. Exacerbations of ILD are rare (0.5%) and only one mortality has been reported (0.2%) (11–20).

Pajares et al conducted a prospective randomised trial comparing TBCLB with TBB and demonstrated a mean specimen size for TBCLB samples of 14.7mm$^2$ (vs 3.3mm$^2$ for TBB biopsy p <0.001) resulting
in a histological diagnosis in 74.4% of patients versus 34.2% in the TBB group (p<0.001) (14). When comparing this technique with surgical lung biopsy there are various potential advantages. General anaesthesia is not necessary and the procedure can be performed as a day case with uncomplicated cases returning home the same day. Future clinical trials and an increase in real world experience of TBCLB is likely to cement it’s use for selected cases, potentially reducing the number of surgical lung biopsies performed.

SURGICAL LUNG BIOPSY

Surgical lung biopsies (SLB) are the current gold standard for obtaining histological material in the diagnosis of clinically and radiologically unclassifiable ILD. SLB is usually performed via the less invasive Video-Assisted Thoracoscopic Surgical (VATS) approach. As previously described the 2011 ATS/ERS consensus statement means about two thirds of IPF cases can be diagnosed on the basis of typical clinical and radiological findings of UIP (Figure e), however, indications are that only 7.5%-12% of suspected IPF patients undergo SLB in the UK (28). This reflects clinicians’ reluctance to refer patients for a procedure associated with a significant mortality and morbidity.

The average hospital stay associated for a VATS biopsy is 2-4 days (23), with mortality rates of 3-4% and an overall complication rate of up to 16% (24). Common complications include; persistent air-leak, exacerbations of underlying ILD due to mechanical stress of single lung ventilation, bleeding and delayed wound healing. In addition, 57% of patients report pain at the incision site 6-12 months after surgery (25). It is also worth remembering that SLB does not guarantee a definite pathological diagnosis, with diagnosis rates ranging from 34% to 100% (23,24).

MULTIDISCIPLINARY TEAM

Taking into consideration the various investigations involved in ILD diagnosis it is clear that no single diagnostic test can provide a confident answer. A consensus approach by a Multidisciplinary Team (MDT) with expertise in ILD is thus considered the gold standard (Figure f). The MDT integrates all available data at several stages of the work-up. This not only improves inter-observer agreement and diagnostic confidence, but may also prevent unnecessary surgical biopsies, whilst identifying patients in whom a biopsy may effectively contribute to the diagnosis (26). Current NICE guidelines recommend that IPF should only be diagnosed by MDT consensus and stipulates a minimum MDT composition (27).
NOVEL THERAPIES IN IPF

There has been a dramatic increase in clinical trial activity in IPF in recent years, with the discovery and approval of two new anti-fibrotic drugs, pirfenidone and nintedanib, heralding a new era in the disease. While these novel anti-fibrotic agents have been shown to slow the decline in Forced Vital Capacity (FVC) they neither halt progression nor reverse existing fibrosis. In part due to considerable cost, their use is restricted by NICE to patients fulfilling certain criteria, namely a FVC of 50-80% predicted, thereby excluding patients at each extreme of the disease process and those with spuriously maintained FVC due to concurrent emphysema. Given these restrictions, as well as the limitations of these therapies, the importance of non-pharmacological therapy such as pulmonary rehabilitation, plus the enrolment of patients into clinical trials (Figure g) should not be underestimated.

NOVEL BIOMARKERS:

The need to distinguish the different IIPs has driven the search for novel diagnostic biomarkers. In addition, there are marked survival differences even within specific groups such as IPF. Biomarkers that can identify these phenotypes are needed for clinical decision-making, but they also have the potential to aid cohort enrichment in clinical trials. Previous landmark studies have beautifully illustrated this need, with variable rates of decline in placebo arms leading to inconsistent results and a delay in drugs being approved (28).

To date, efforts have focussed on serum biomarkers (29) that are relatively easy to access and novel imaging modalities which potentially inform on disease activity within the lung. In particular, Positron emission tomography (PET) allows non-invasive measurement of cellular metabolism in vivo. The 18F-Fluorodeoxyglucose (18F-FDG) PET signal is consistently raised in ILD (30) and is both stable and reproducible (Figure h). In a population of over 200 ILD patients, we have shown that baseline measures of 18F-FDG up-take on PET are related to patient survival in both IPF and other IIPs (paper in preparation). High pulmonary 18F-FDG up-take is associated with poor survival, giving additional information to pulmonary function testing (PFT) and thus, could become a valuable part of the initial work up in newly diagnosed patients.

CONCLUSION

The diagnosis of ILD is a challenging and involved process. It relies on detailed history taking and the integration of various investigations and specialities. The relative rarity of these diseases makes distinguishing subtypes even more difficult for clinicians with a mixed respiratory case workload and
thus, infrequent exposure to ILD. Having said this, the incidence of ILD is increasing and there is potential for specialist centres to become overwhelmed with patients, putting a greater than ever emphasis on collaboration with referring centres and a concerted effort to employ a hub and spoke model. This has the added advantage of facilitating a more patient focussed approach, minimising the need for unnecessary travel and facilitating access to ancillary local services, such as pulmonary rehab, oxygen providers and palliative care services.

Irrespective of expertise, uncertainty is inherent in the diagnosis of these diseases, although arguably encountering ILD on a frequent basis allows the physician to become more comfortable with these uncertainties, thus embracing the concept of continuous diagnostic review. The hope remains that in time, reliable, non-invasive biomarkers will identify disease subtypes, predict prognosis and potentially replace the need for biopsy. Much of the heterogeneity seen in IPF may be explained by the existence of endotypes, in other words, mechanistically different disease subtypes, which consequently exhibit very different responses to therapy. Future treatments therefore have the potential to be greatly influenced by identifying these groups through the use of genetic testing and a move towards personalised disease management.

References;


Figure a: Schematic classification of Interstitial Lung Diseases according to aetiology. The finding of histological UIP in a patient with an IIP leads to the specific diagnosis of Idiopathic Pulmonary Fibrosis (IPF).
Figure b: The Pathogenesis of IPF. In an initiating phase, there is lung alveolar epithelial damage with loss of the normal lung architecture and disruption of the basement membrane across which gas exchange takes place. With further epithelial damage and apoptosis, comes up-regulation of epithelial integrins such as αvβ6 and a phase of fibroproliferative repair dominates driven by high levels of TGFβ. Released in an inactive form, this cytokine requires an activation step facilitated by integrins that bind the RGD motif of pro-TGFβ and promote its cleavage and activation. Locally activated TGFβ drives the recruitment of fibroblasts and a feed-forward cycle of further TGFβ production. Under these conditions, fibroblasts differentiate into myofibroblasts that express high levels of integrin αvβ6, are resistant to apoptosis and lay down collagen matrix. Once collagen has been laid down in a lung, the architecture of which is already distorted, gas exchange is no longer efficient. There is a change in the vasculature of the lung parenchyma with both fall-out of blood vessels and neo-angiogenesis driven by local production of vascular endothelial and platelet derived growth factors (VEGF and PDGF). At this final phase the lung is irreversibly scarred.

34x25mm (600 x 600 DPI)
Figure c: Diagnostic criteria for a definite UIP pattern on HRCT:
1. subpleural, basal predominance; (red).
2. reticular abnormality (blue).
3. honeycombing with or without traction bronchiectasis (yellow).
4. absence of features inconsistent with UIP pattern

Figure c
Figure d: Comparison between transbronchial cryo lung biopsy on the left and traditional forceps transbronchial biopsy on the right performed on the same patient at two different sittings. TBCLB shows preserved architecture of parenchymal tissue and a total biopsy area of 46.81 mm² and mean biopsy area of 11.7 mm². TBBx is characterised by crush and haemorrhagic artefact and a total biopsy area of 14.11 mm² and mean biopsy area of 2.8 mm².
Figure e: Diagnostic algorithm for idiopathic pulmonary fibrosis, in part adapted from the ATS/ERS consensus statement
Figure f: The role of Specialist MDTs and Specialist Referral Centres in the diagnosis and management of ILD.

Referring Centre assesses patient including discussion at local MDT

- ILD – does not fit any pattern or rare ie LAM, LCH
- Definite or Probable IPF
- Definite or possible NSIP +/- CTD
- Patient request or to participate in clinical trial
- Lung transplantation being considered

Review at specialist centre MDT

- Suitable for anti-fibrotic treatment or possible clinical trial
- For diagnostic review including possible biopsy
- For diagnostic review including possible biopsy
- Progressive in spite of treatment
- Unable to give desired treatment i.e. cyclophosphamide locally

Feedback provided and initiated locally

Review at Specialist Centre with shared care/shared follow up approach encouraged

https://mc04.manuscriptcentral.com/clinmed
Figure g: Schematic ILD treatment algorithm. *No robust evidence for managing exacerbations with variation between centres, should be discussed with specialist centre if possible. GOR; Gastro-oesophageal reflux. PHT; pulmonary hypertension

Figure g
Figure h: PET signal superimposed on HRCT scan of a patient with UIP. A region of interest has been drawn around the area of highest SUV.
<table>
<thead>
<tr>
<th>Antibody</th>
<th>Associated CTD</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANA (&gt;1:320)</td>
<td>Many</td>
</tr>
<tr>
<td>RF (&gt;60 IU/mL)</td>
<td>RA, Sjögrens, SLE</td>
</tr>
<tr>
<td>Anti-CCP</td>
<td>RA</td>
</tr>
<tr>
<td>Anti-centromere</td>
<td>Systemic sclerosis</td>
</tr>
<tr>
<td>Anti-nuclear ANA</td>
<td>Systemic sclerosis</td>
</tr>
<tr>
<td>Anti Ro (SS-A)</td>
<td>SLE, Sjögrens and others</td>
</tr>
<tr>
<td>Anti La (SS-B)</td>
<td>SLE, Sjögrens</td>
</tr>
<tr>
<td>Anti-RNP</td>
<td>SLE, MCTD</td>
</tr>
<tr>
<td>Anti-dsDNA</td>
<td>SLE</td>
</tr>
<tr>
<td>Anti-Smith</td>
<td>SLE</td>
</tr>
<tr>
<td>Anti tRNA synthetase</td>
<td>Poly-/dermatomyositis (anti-synthetase syndrome)</td>
</tr>
<tr>
<td>Anti-PM-Scl</td>
<td>Systemic sclerosis/ myositis overlap</td>
</tr>
<tr>
<td>Anti-Th/To</td>
<td>Systemic sclerosis</td>
</tr>
<tr>
<td>Anti-U3 RNP</td>
<td>Systemic sclerosis</td>
</tr>
<tr>
<td>ANCA panel</td>
<td>Vasculitides</td>
</tr>
<tr>
<td>Anti topoisomerase (Scl-70)</td>
<td>Systemic sclerosis</td>
</tr>
</tbody>
</table>

Table 1: Autoantibodies in Connective Tissue ILDs

ANA: antinuclear antibody; RF: Rheumatoid Factor; Anti-CCP: anti-cyclic citrullinated peptide antibody anti-RNP: anti-Ribonucleoprotein; anti-dsDNA: anti-double stranded DNA; Anti-PM-Scl: anti polymyositis-scleroderma; ANCA: Anti-neutrophil cytoplasmic antibodies; RA: rheumatoid arthritis; SLE: Systemic lupus erythematosus; MCTD: mixed connective-tissue disease
Optimal HRCT technique for evaluation of ILD

• Non-Contrast scans obtained on full inspiration without respiratory motion
• Contiguous or non-contiguous axial scans with thin sections, reconstructed at ≤2 cm intervals
• Reconstructed slice collimation ≤2 mm
• High resolution reconstruction algorithm
• Field of view to include lungs only
• Expiratory scans are helpful to exclude lobular air trapping suggestive of hypersensitivity pneumonitis
• Prone scans if dependent density obscures detail on supine images
• Optional coronal and sagittal reconstructions if volumetric images are obtained

Table 2: The ATS/ERS consensus statement for the diagnosis of IPF set out criteria for the optimal HRCT technique for evaluation of ILD