Review

The utility of comprehensive autoantibody testing to differentiate connective tissue disease associated and idiopathic interstitial lung disease subgroup cases

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Abstract

Interstitial lung disease (ILD) comprises many heterogeneous disease groups, the largest being CTD-associated and those labelled as idiopathic out of necessity. The mechanisms causing ILD are poorly understood, but most CTD- and idiopathic-ILD cases can respond to immunosuppression, clearly suggesting a pathological role for inflammation. By contrast, corticosteroid immunosuppression causes harm without benefit in the feared idiopathic pulmonary fibrosis, suggesting that inflammation plays little pathological role, and where ILD progresses rapidly to lethal outcome even with anti-fibrotic drug use. Given the treatment response differences apparent between ILD subgroups, and the dangers and costs of corticosteroid and anti-fibrotic drug use, respectively, it has become vital in every ILD patient to make an accurate subgroup diagnosis, to optimize treatment selections. This review discusses why differentiating CTD- and idiopathic-ILD subgroup cases remains so problematic, and why existing comprehensive CTD-specific serology would, if generally available, represent an ideal biomarker tool to enhance ILD subgroup diagnostic accuracy.

Key words: interstitial lung disease, connective tissue disease, antibodies, biomarkers, serology, myositis, anti-synthetase syndrome.

Introduction

The parenchymal tissues of the terminal airways comprise the alveolar epithelium and basement membrane, the peri-vascular/peri-lymphatic (i.e. the true) interstitial space and the capillary basement membrane and epithelium. Damage to any of these tissues could lead to diffuse parenchymal lung disease, although interstitial lung disease (ILD) is the preferred UK term [1]. The diffuse parenchymal lung disease pattern distinguishes ILD from other lung pathologies, including those affecting the pleura and larger airways. If ILD-associated parenchymal injury, and any associated inflammation, is not arrested to facilitate
healing, irreversible pulmonary scarring and fibrosis may follow [2]. It is the combination of parenchymal injury, primary and/or secondary inflammation, pulmonary fibrosis and the resulting dyspnoea that physicians recognize overall as ILD. Impaired gas transfer can occur early, if inflammatory infiltrations are sufficiently severe, or only once fibrosis has become substantial, irrespective of cause. In individual cases the extent of parenchymal inflammation, relative to that of fibrosis, will depend on the mechanism causing injury, and also dictate the effectiveness of immunosuppressive therapies [3]. Patients accurately diagnosed and treated will normally fare better. Non-responsive or untreated patients usually progress from being dyspnoeic on exertion only to being dyspnoeic even at rest. Fatal outcomes ensue from respiratory failure, or its cardiac complications [4].

Why the difficulties in diagnosing CTD-associated ILD?

A representative classification of ILDs is shown in Fig. 1 [5]. Though apparently simple, this classification hides many potential pitfalls. The two largest ILD groups are the CTD-associated ILDs (CTD-ILDs) and the idiopathic-ILDs (also known as the idiopathic interstitial pneumonias, or IIPs). Both groups have many subgroups. Differentiating between subgroup cases can prove considerably difficult because their ILD features can overlap, as can be demonstrated on lung biopsy material where available, and on high resolution CT (HRCT) scans [6]. Non-specific interstitial pneumonia (NSIP) and usual interstitial pneumonia (UIP) patterns on HRCT can be seen in CTD-ILD and idiopathic-ILD [7]. The NSIP pattern is characterized by ground glass changes typically present at the periphery and bases of the lungs, whereas a UIP pattern shows peripherally distributed reticular and fibrotic changes typically affecting the lung bases with honeycomb features, and the absence of significant ground glass changes [8]. The commonest ILD is idiopathic pulmonary fibrosis (IPF), where scans typically demonstrate the UIP pattern (IPF and idiopathic UIP are thus interchangeable terms). IPF/idiopathic UIP is the most feared of all ILDs, because it is so relatively common and is associated with a ∼50% mortality within only 3 years of ILD onset despite all treatments [4] (see later). When a physician accurately diagnoses a CTD in a patient with NSIP, that case would obviously be labelled as a CTD-ILD, and the detected CTD is assumed to be the underlying cause of the NSIP. If, on the other hand, no CTD signs were detectable, and no other cause for an NSIP pattern was apparent, that case would of necessity be assigned an idiopathic-ILD label, that is, idiopathic NSIP. In each instance, the final ILD diagnosis will have relied on use of the combined clinical, radiological, serological and (where available) lung histological features. However, this seemingly logical process can break down with CTDs. This is because ILD may be the presenting sign of a CTD, that is, where extra-pulmonary CTD features have yet to appear [9–11]. ILD thus represents a forme fruste CTD. Furthermore, CTD symptoms may be present but of only very subtle character, and so easily missed in a busy respiratory clinic. For example, in the anti-synthetase syndrome the classic CTD features other than ILD (i.e. Raynaud’s

**Fig. 1** A classification of ILD into groups and subgroups

![Interstitial Lung Disease](https://academic.oup.com/rheumatology/article-abstract/56/8/1264/2433519)

The IPF and RA-ILD (UIP) subgroups are formatted left in each box, and made bold, to highlight that these ILDs exhibit similar fibrotic HRCT patterns (UIP) and are similarly treatment-resistant, with rapid progression to lethal outcomes within only 3–5 years of ILD onset. In contrast, and formatted to the right in each box, the other IIP and CTD-ILD subgroups, the latter including RA-ILD (non-UIP), show varying degrees of cellularity (i.e ground glass) on HRCT, and are usually responsive to immunosuppression to some extent. IPF: Idiopathic Pulmonary Fibrosis, ILD: Interstitial Lung Disease, UIP: Usual Interstitial Pneumonia, HRCT: High Resolution Computed Tomography, IIP: Idiopathic Interstitial Pneumonia. (Adapted from Ryerson CJ, Collard HR. Update on the diagnosis and classification of ILD. Curr Opin Pulm Med 2013;19:453–9).
 syndrome, myositis, mechanic’s hands, fever, rash, inflammatory arthritis) may be absent or very subtle only [12]. Moreover, if ILD is successfully treated early, the immunosuppressive agents used may well preclude the development of CTD signs other than ILD. Physicians should employ serology to screen for a CTD in every ILD case, though chest physicians may employ serology less rigorously than would perhaps be expected of a rheumatologist [13]. False positive serology and the existence of sero-negative CTDs would also cause difficulties. A low suspicion for the presence of a CTD may mean that some CTD-ILD patients never see a rheumatologist.

The diagnostic limitations of HRCT

HRCT is an important assessment tool to differentiate between fibrosis and inflammation in ILD patients. Where clinical history and examination fail to confirm a clear ILD diagnosis, and no CTD symptoms or signs are apparent, clinicians must then rely on HRCT, if serology is diagnosis inadequately or unhelpful. However, HRCT has poor utility as a stand-alone diagnostic tool. For instance, HRCT appearances have no proven associations with any defined disease processes. Also, not only do the same HRCT patterns occur in different ILD subgroups, but different HRCT patterns can occur in the same disease subgroup [14]. Furthermore, similar HRCT patterns can be associated with differential outcomes. For example, the most common HRCT appearances in SSC are those of NSIP or UIP, the latter thus mimicking IPF. However, in contrast to IPF, SSC-ILD patients with UIP can respond to immunosuppression [15–17]. Honeycombing is a recognized HRCT feature of IPF, but honeycombing can also occur in myositis-ILD, thus again mimicking IPF, yet myositis-UIP cases can respond to immunosuppression [18]. In the absence of comprehensive CTD serology, inaccurate diagnostic assignments by HRCT will have occurred previously, and likely will continue to occur. That fibrotic HRCT changes respond differently in IPF than in CTD-ILDs with predominantly fibrotic HRCT changes may reflect differential disease mechanisms. Such a range and variability of HRCT appearances, clinical associations and treatment responses highlights the problems of diagnostic uncertainty when CTD-specific serology is inadequate. In addition, there are the difficulties that HRCT patterns may overlap, and that inter-observer variability of image analysis also occurs [19, 20].

The 2011 American Thoracic Society (ATS)/European Respiratory Society (ERS) guidelines for the diagnosis and management of IPF suggested that an accurate ILD diagnosis is optimally secured by the combination of a thorough medical history (to detect causative CTDs or occupational or environmental exposures, etc.), a careful clinical examination (to detect CTD or granulomatous signs, etc.), serological screening, an HRCT scan and lung histology where feasible, though co-morbidities dictate that only a minority of UK cases are sufficiently fit for lung biopsy [21]. A definite occupational, iatrogenic, granulomatous, CTD, etc. history and/or detection of clinical signs would likely secure an underlying ILD diagnosis, and so preclude the need for lung biopsy. However, when diagnostic doubts remain, final diagnostic decisions are increasingly made in the UK by expert multidisciplinary teams in tertiary ILD clinics, with input from pulmonary clinicians, radiologists, pathologists and increasingly rheumatologists with an interest in ILD. Tertiary ILD service developments have in part been driven by the advent of newer drugs for IPF, a particularly challenging ILD that is preceded by little or no inflammation, and that is therefore non-responsive to immunosuppression [22, 23].

The diagnostic use of serology in the absence of ILD-specific biomarkers

A factor that has to date critically limited ILD subgroup diagnostic capability has been the lack of reliable, ILD-specific biomarkers. UK physicians, including rheumatologists, have thus struggled diagnostically with idiopathic ILD and with CTD-ILD when CTD signs other than ILD are absent, though this situation is set to improve. IPF can only be diagnosed once a CTD has been definitively excluded, but the 2011 and 2013 ATS/ERS guidelines gave only very limited advice regarding the stringency of serology required to interrogate for a CTD [21, 24]. Thus, and even in most tertiary UK ILD clinics, CTD exclusion continues to rely on routine immunology, that is, immunology testing for rheumatoid factor, anti-CCP antibodies, ANA and extractable nuclear antigens, the latter usually limited to anti-Ro/La/Jo/Sm/RNP/Scl-70. A recent ERS/ATS research statement on interstitial pneumonia with autoimmune features suggests, for the first time, that classification criteria should contain a serological domain [25]. This statement suggests use of serology more extensive than previously advised in any ATS/ERS guidelines [8, 21, 24], though no details are given regarding how to use this newly advocated serology, or regarding its diagnostic utility [25]. Moreover, for many of the newer antibody specificities listed in this update, and especially for some of those associated with myositis, serological detection systems are unfortunately not currently available in routine UK clinical practice. Thus, in myositis-ILD, a growing number of myositis-specific autoantibodies (MSAs) and myositis-associated autoantibodies (MAAs) are now detectable (Tables 1 and 2) [26]. MSA/MAAs accurately predict the presence of myositis clinical features, including the likelihood of developing an associated ILD, so these antibodies represent surrogate ILD biomarkers, and hence their inclusion in the most recent ERS/ATS update [25]. Currently, however, many MSA/MAAs are only detectable by expensive immunoprecipitation techniques [26], although immunoblot technology for a number of these antibodies is now available in some National Health Service hospitals. Even when these are available, issues of diagnostic accuracy and nomenclature still arise. For instance, in the case of anti-PL-7 and anti-PL-12, these antibodies often associate with ILD in the absence of myositis [11, 27], that is, in amyopathic ILD. Affected patients may not initially or ever exhibit CTD features other than their ILD [11, 12]. Reliable detection of an

Autoantibody. (Table adapted from Betteridge ZE et al. Novel autoantibodies and clinical phenotypes in adult and juvenile myositis. Arthritis Res Ther 2011;13:209.)

Table 1: Clinical associations of the known myositis-specific autoantibodies and the likelihood of antibody-positive patients developing an associated interstitial lung disease

<table>
<thead>
<tr>
<th>MSA</th>
<th>Prevalence in myositis (%)</th>
<th>Clinical associations</th>
<th>Likelihood of developing ILD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-synthetases:</td>
<td></td>
<td>Associated with anti-synthetase syndrome, and characterized by the presence of:</td>
<td>~70% develop ILD [30]</td>
</tr>
<tr>
<td>Anti-Jo-1 (histidyl)</td>
<td>15-20</td>
<td>Myositis (PM or DM)</td>
<td>~90% develop ILD, myositis often mild [11]</td>
</tr>
<tr>
<td>Anti-PL-12 (alanyl)</td>
<td>&lt;5</td>
<td>Arthritis ILD</td>
<td>~90% develop ILD, often prior to myositis [31]</td>
</tr>
<tr>
<td>Anti-PL-7 (threonyl)</td>
<td>&lt;5</td>
<td>Raynaud’s Fevers</td>
<td></td>
</tr>
<tr>
<td>Anti-KS (asparaginyl)</td>
<td>&lt;5</td>
<td>Mechanic’s hands</td>
<td></td>
</tr>
<tr>
<td>Anti-OJ (isoleucyl)</td>
<td>&lt;5</td>
<td>Rapidly progressive ILD, especially in Japanese and Chinese patients</td>
<td>50-70% develop rapidly progressive ILD in Japanese/Chinese ethnicity [36, 37], ~50% in Caucasian [38]</td>
</tr>
<tr>
<td>Anti-EJ (glycyl)</td>
<td>&lt;5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anti-Zo (phenylalanyl)</td>
<td>&lt;1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anti-Ha (tyrosyl)</td>
<td>&lt;1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anti-MDA5</td>
<td>Associated with clinically amyopathic DM</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The following MSAs are rarely if ever associated with likelihood of developing an ILD: anti-Mi-2, anti-NXP2, anti-TIF1-γ, anti-SAE and anti-SRP. ILD: interstitial lung disease; MDA5: melanoma differentiation-associated gene 5; MSA: myositis-specific autoantibody. (Table adapted from Betteridge ZE et al. Novel autoantibodies and clinical phenotypes in adult and juvenile myositis. Arthritis Res Ther 2011;13:209.)

ILD-only MSA/MAA here would presumably secure a CTD-ILD diagnosis, and it could be argued that detected ILD-only MSA/MAAs could be appropriately renamed as ILD-specific/associated autoantibodies (i.e. ISA/IAA), or as CTD-ILD-specific/associated autoantibodies, though the latter appears somewhat cumbersome, especially regarding the issue of an acronym. As the intracellular targets of many MSA/MAAs, and of all eight of the known anti-synthetases, are cytoplasmic rather than nuclear, a negative ANA on routine screening does not of itself exclude a CTD-ILD diagnosis. However, the staining patterns observed when ANA screening is undertaken by indirect immunofluorescence testing on HEp-2 cells will potentially disclose the presence of a CTD [28, 29]. The HEp-2 technique is rapid and inexpensive, so if available its use for screening ILD patients for an underlying CTD is recommended.

The recently described anti-melanoma differentiation-associated gene 5 (MDA5) antibody occurs in DM patients who are clinically amyopathic, but who may suffer a very aggressive ILD, that is, one proving lethal within months or only weeks of ILD onset, despite all immunosuppressive interventions [36]. Anti-MDA5 could thus also be termed an ISA. Fortunately this antibody and its associated ILD are rare in the UK [26]. In other CTDs with a potential to develop ILD with or without myositis, for example, in mixed CTD, many patients will also possess one or more MAAs. In SSc there are many SSc-specific antibodies that are also strongly associated with ILD development (see Table 2). Thus, MSAs, MAAs and SSc-associated antibodies should all be regarded as surrogate biomarkers of ILD, or when appropriate even as ISA/IAAs. The recent ERS/ATS update recognizes the growing evidence suggesting that serology should be utilized more readily to enhance CTD-ILD diagnosis. However, this update also points out that detailed prospective studies are now required to validate the newly proposed classification criteria for interstitial pneumonia with autoimmune features [25]. In view of the current inadequacies of routine serology, and given that respiratory physicians may lack expertise in assessing extrapulmonary CTD clinical features [25], some CTD patients with HRCT appearances suggestive of UIP will be misdiagnosed as IPF. Similarly, some patients may be assigned an idiopathic-ILD label when they actually have a CTD-ILD that remains immunologically undisclosed.

Differential outcomes in CTD-ILDs

In the past, CTD- and idiopathic-ILD patients with similar HRCT patterns would have been treated with similar immunosuppressive regimes, so diagnostic labels then were therapeutically of lesser importance. However, recent mechanistic research and clinical trials in IPF have dramatically altered the treatment landscape. Making an accurate ILD diagnosis, to differentiate a CTD-ILD from an idiopathic-ILD, has thus become crucial, as treatments may differ markedly, and especially between IPF and most other ILD subgroups. Following the publication of the PANTHER study, which examined the impact of
### Table 2: Clinical associations of various SSc-associated antibodies and myositis-associated antibodies and the likelihood of antibody-positive patients developing interstitial lung disease

<table>
<thead>
<tr>
<th>SSC-associated antibody or MAA</th>
<th>Prevalence of antibody</th>
<th>Disease associations</th>
<th>Likelihood of developing ILD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-topoisomerase (ScI70)</td>
<td>~30% of SSc patients [39]</td>
<td>dcSSc [40]</td>
<td>~60% of patients develop ILD [41]</td>
</tr>
<tr>
<td>Anti-Th/To</td>
<td>~5% of SSc patients [40]</td>
<td>lcSSc [42]</td>
<td>~50% of patients develop ILD [43]</td>
</tr>
<tr>
<td>Anti-U3 RNP</td>
<td>~8% of patients with SSc [44]</td>
<td>Occurs in SSc. Associated with skeletal muscle involvement and pulmonary arterial hypertension [45]</td>
<td>~40% of patients have ILD [45]</td>
</tr>
<tr>
<td>Anti-U11/U12 RNP</td>
<td>~3% of patients with SSc [46]</td>
<td>Occurs in SSc</td>
<td>~80% of patients develop ILD which is often severe [46]</td>
</tr>
<tr>
<td>Anti-RuvBL1/2</td>
<td>1-2% of SSc patients [47]</td>
<td>Associated with myositis overlap and diffuse skin thickening [47]</td>
<td>Over 50% of patients will develop ILD [47]</td>
</tr>
<tr>
<td>Anti-Eif2B</td>
<td>~1% of patients with SSc/ SSc overlap [48]</td>
<td>Associated with SSc/SSc overlap syndrome</td>
<td>Up to 100% of patients will develop ILD [48]</td>
</tr>
<tr>
<td>Anti-PM-Scl</td>
<td>Occurs in up to 17% of patients with overlap myositis [49], 3-6% of patients with SSc [50]</td>
<td>Associated with scleromyositis</td>
<td>~50% of patients will develop ILD [49, 51]</td>
</tr>
<tr>
<td>Anti-Ku</td>
<td>Occurs in 13% of patients with overlap myositis [49]</td>
<td>Associated with myositis overlap syndrome</td>
<td>~30% of patients will develop ILD [49]</td>
</tr>
<tr>
<td>Anti-U1 RNP</td>
<td>5-35% of patients presenting with SSc or overlap syndromes [50]</td>
<td>Associated with MCTD</td>
<td>~35% show HRCT abnormalities associated with ILD, ~20% classified as severe [52]</td>
</tr>
<tr>
<td>Anti-Ro-52/60</td>
<td>~40% of IIM patients have anti-Ro-52/60 alongside their MSA or MAA [51]</td>
<td>Frequently occur with an MSA, especially with the anti-synthetases, or with various MAAs</td>
<td>~40% of patients develop ILD, however it is unlikely that these antibodies are responsible for the ILD risk, which is instead due to the MSA/MAA detected [51]</td>
</tr>
</tbody>
</table>

The following SSc-associated antibodies are less frequently associated with ILD: anti-centromere antibody and anti-RNA polymerase III. Table compiled from the cited references. HRCT: high resolution CT; IIM: idiopathic inflammatory myopathy; ILD: interstitial lung disease; MAA: myositis-associated antibody; MCTD, mixed connective tissue disease; MSA: myositis-specific antibody.

Treating IPF patients with high-dose prednisolone co-prescribed with AZA, this immuno-suppressive regime was deemed contra-indicated because of its thus proven iatrogenic dangers [23]. New anti-fibrotic agents such as pirfenidone and nintedanib are now licensed and available for use to slow IPF disease progression [53–55]. These agents are, however, considerably expensive, so their use in England can only be recommended when an IPF diagnosis is deemed robust, that is, when made in a designated tertiary ILD centre, and according to the 2011–13 ATS/ERS guidelines [21, 24].

Most CTD-ILD cases have a capacity to respond to immunosuppression. If a degree of fibrosis has already occurred, suppressing residual inflammation would likely still act to limit fibrotic progression, thus stabilizing dyspnoea and associated disability. Such treatment responsiveness supports the notion that in most CTD-ILD cases, including myositis-ILD, ground glass changes reflect a suppressible inflammation component [56]. That treatment responses are also seen where fibrotic HRCT changes predominate in some CTD-ILD cases is poorly understood. A generic and suppressible inflammatory component could be assumed in all CTD-ILD cases, yet treatment outcomes are not always good. For instance, when RA patients develop an associated ILD (RA-ILD) with a UIP pattern on HRCT, that is, RA-ILD (UIP), such cases are notoriously non-responsive to immunosuppression [57], RA-ILD (UIP) is associated with a lethal outcome within only 3 years of ILD onset, thus clearly mimicking IPF [57, 58]. Moreover, the RA-ILD (UIP) pattern on HRCT is identical to that of IPF. Therefore, although RA-ILD (UIP) is much rarer than IPF, it is a diagnosis just as feared as IPF. No research to date has reported on whether anti-fibrotic agents have a therapeutic role to play in RA-ILD (UIP). Some RA-ILD patients may develop organising pneumonia within only 3 years of ILD onset, that is, RA-ILD (UIP). In this context, HRCT changes predominate in some CTD-ILD cases is poorly understood. A generic and suppressible inflammatory component could be assumed in all CTD-ILD cases, yet treatment outcomes are not always good. For instance, when RA patients develop an associated ILD (RA-ILD) with a UIP pattern on HRCT, that is, RA-ILD (UIP), such cases are notoriously non-responsive to immunosuppression [57], RA-ILD (UIP) is associated with a lethal outcome within only 3 years of ILD onset, thus clearly mimicking IPF [57, 58]. RA-ILD (UIP) is associated with a lethal outcome within only 3 years of ILD onset, thus clearly mimicking IPF [57, 58].
Academic issues

Given that the 2011 ATS/ERS guidelines for IPF [21] were constructed without reference to comprehensive serology to test for MSA or MAA and SSc-associated autoantibodies, a question naturally arises regarding the robustness of an IPF diagnosis when made without such serology. Such a stricture may well have blighted previous mechanistic IPF research. For instance, in the few research studies in which relatively comprehensive myositis serology testing was undertaken in IPF cases, who would by definition have required IPF-consistent HRCT changes for study inclusion (i.e. UIP, or fibrotic NSIP), the results demonstrated such serology to be positive in a significant number of cases [9, 59]. As MSAs have highly significant HLA associations [60], these results could imply that IPF also has strong HLA associations, yet genetic studies using genome-wide association scan technology in IPF have failed to demonstrate significant HLA associations [61, 62]. Moreover, many of the MSA-positive cases in these studies were younger females, an observation more in keeping with a CTD-ILD rather than an IPF phenotype. Some of the MSA-positive IPF cases were presumably myositis-ILD cases, but where the CTD diagnosis had remained covert until the research myositis serology was undertaken. In contrast, in a Mexican study, highly significant HLA associations were found in IPF [63]. In this study the IPF diagnoses were based on the 2002 ATS/ERS IIP diagnostic guidelines, which gave no guidance on use of serology to interrogate for the presence of a CTD [8]. The highly significant HLA associations detected in this study, with odds ratios >10, could again imply that their IPF case cohort was in fact contaminated by many covert CTD-ILD cases, such as those with MSA/MAA. The contradictory nature of these genetic results suggests that, even in a phenotype as apparently robust as IPF diagnosed strictly in a specialized ILD clinic setting, case stratification errors have likely still occurred. To optimize accuracy of case stratifications to guarantee homogeneous cohorts for future IPF genetic studies will clearly require modern and comprehensive serology to be used to definitively exclude all CTD cases. Amyopathic ILD patients without a rash, and especially those with an anti-synthetase other than anti-Jo-1, would clearly cause confusion in any ILD study where comprehensive serology was not available. If all patients diagnosed as IPF in a tertiary UK ILD clinic were interrogated by immunoprecipitation, would a substantial cohort prove positive for CTD-associated antibodies? If so, then undertaking comprehensive serology on all incident ILD cases without obvious CTD signs would seem justifiable, at least until this question has been addressed.

Conclusions

The relative paucity of aetopathological insights so far gained in ILD reflects the difficulties of accurately assigning ILD subgroup diagnoses without comprehensive CTD serology, and that case-stratification errors have likely occurred in previous research. This may have contributed to the apparent confusion in the literature regarding diagnostic labels, mechanistic issues and the apparent contradictory nature of genetic research outputs. These insight deficits also reflect the rarity of ILD, and the invasiveness of lung biopsy procedures has severely hampered investigations of the ILD organ target. To facilitate accurate future clinical ILD care, so as to optimize outcomes, it will be vital that comprehensive CTD-serotyping becomes standard in routine practice, though substantial technological developments will be required to achieve this. Until then, serology by immunoprecipitation will continue to be crucial, and especially in the ILD research setting. Given the extreme rarity of some ILD subgroups, and the sample size problems that would arise for instance during between-subgroup comparative research, it is essential that multicentre collaborative efforts develop to recruit the large ILD subgroup cohorts required to facilitate such research. Such cohorts would make it possible to prospectively correlate ILD clinical phenotypes with serotypes and HRCT-generated radiological phenotypes for all ILD subgroups. Examining the validity of HRCT to assign idiopathic-ILD subgroup diagnoses would also become possible. It is feasible that some idiopathic subgroups, such as idiopathic NSIP and COP, will become smaller or even disappear, as more individual idiopathic-ILD cases are appropriately reassigned into CTD-ILD subgroups. Serologically better defined ILD subgroups will ensure more meaningful between-subgroup evaluations, for instance in future immunogenetic comparisons. The use of larger, more homogeneous subgroup cohorts will clearly facilitate ILD research, and thus ultimately improve quality of patient care.

Acknowledgements

The authors of this review are co-ordinating a UK-wide multicentre collaborative recruitment of ILD cases, initially called: identifying disease susceptibility genes and autoantibodies associated with the development and clinical characteristics of interstitial lung disease (ILD) in patients with and without proven connective tissue diseases (CTDs), Version 1.0 18 December 2010. This was subsequently renamed: the UK Biomarkers in Interstitial Lung Disease (UK-BILD) Study. This is NIHR-approved (UKCRN ID: 15775), so case recruitment is eligible for CRN network support (http://public.ukcrn.org.uk/search/). Centres interested in participating in UK-BILD should contact Mr Paul New, who will facilitate the local ethical approval process (paul.new@liverpool.ac.uk). CC received PhD funding from the the Institute of Ageing and Chronic Disease as well as Arrowe Park Endowment Funds.

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