Position Paper

Interstitial lung abnormalities detected incidentally on CT: a Position Paper from the Fleischner Society

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The term interstitial lung abnormalities refers to specific CT findings that are potentially compatible with interstitial lung disease in patients without clinical suspicion of the disease. Interstitial lung abnormalities are increasingly recognised as a common feature on CT of the lung in older individuals, occurring in 4–9% of smokers and 2–7% of non-smokers. Identification of interstitial lung abnormalities will increase with implementation of lung cancer screening, along with increased use of CT for other diagnostic purposes. These abnormalities are associated with radiological progression, increased mortality, and the risk of complications from medical interventions, such as chemotherapy and surgery. Management requires distinguishing interstitial lung abnormalities that represent clinically significant interstitial lung disease from those that are subclinical. In particular, it is important to identify the subpleural fibrotic subtype, which is more likely to progress and to be associated with mortality. This multidisciplinary Position Paper by the Fleischner Society addresses important issues regarding interstitial lung abnormalities, including standardisation of the definition and terminology; predisposing risk factors; clinical outcomes; options for initial evaluation, monitoring, and management; the role of quantitative evaluation; and future research needs.

Introduction

Interstitial lung disease (ILD) comprises a diverse group of lung diseases with overlapping clinical, radiological, physiological, and pathological features.1 Interstitial lung abnormalities (ILAs) refer to the presence of CT scan findings that are potentially compatible with ILD in patients who have partial (eg, abdominal CT including the lower lung zones) or complete chest CT examinations without previous clinical suspicion of ILD. As ILAs are associated with respiratory symptoms, functional impairment, risk of progression, and increased all-cause mortality,2–10 their identification has clinical implications. The term ILAs does not imply the absence of respiratory signs, symptoms, or functional impairment, but when these clinically significant findings are present, ILAs are likely to represent mild ILD rather than subclinical abnormalities. The definition of ILAs is purely radiological and is based on the incidental identification of CT abnormality. Differentiation between ILAs and clinical and subclinical ILD must be on the basis of clinical evaluation.

ILAs are increasingly recognised on chest CT scans.2 Systematic evaluation of large cohorts has shown a prevalence of ILAs in older individuals (>60 years) of 4–9% in smokers and 2–7% in non-smokers (table).11–13 However, their presence is not routinely recorded on radiology reports, even at academic centres.14 ILAs are likely to be increasingly identified with the implementation of lung cancer screening and increased use of CT for other diagnostic purposes. Still, our understanding of ILAs is minimal, with insufficient evidence to provide definitive management recommendations. This Fleischner Society Position Paper provides multidisciplinary perspectives on definition and terminology; predisposing risk factors; clinical outcomes; options for initial evaluation, monitoring, and management; the role of quantitative evaluation; and future research needs.

What definition and terminology could be used to describe and characterise ILAs?

High-resolution CT is highly sensitive for detecting subclinical interstitial abnormalities in high-risk populations, such as patients with connective tissue disease (eg, systemic sclerosis) or occupational exposures (eg, asbestosis).15-21 Systematic evaluation of large cohorts of smokers screened by CT for lung cancer, or undergoing CT as part of epidemiological evaluation of chronic obstructive pulmonary disease (COPD) or cardiovascular risk factors, has shown that these incidental abnormalities are relatively common, particularly in older individuals (table).22-25 Terms applied to this finding have included
interstitial lung changes at an early phase, early ILD,\(^7\) subclinical ILD,\(^6\) and preclinical ILD.\(^8\) Quantitative abnormalities, such as an abnormally high proportion of high-attenuation areas of the lung, have also been identified in cohort studies and are thought to suggest subclinical parenchymal lung disease.\(^9\)

ILAs are not synonymous with subclinical ILD because a subset of patients with ILAs has symptoms and lung function impairment without suspected ILD. A further subset of patients with ILAs is at risk of progression to clinically significant disease. Abnormalities identified during screening for ILD in high-risk groups (eg, those with rheumatoid arthritis, systemic sclerosis, or familial ILD) are not considered as ILAs because they are not incidental; these might be referred to as preclinical ILD and their management is beyond the scope of this Position Paper.

ILAs have been described as non-dependent abnormalities affecting more than 5% of any lung zone (upper, middle, and lower lung zones are demarcated by the levels of the inferior aortic arch and right inferior pulmonary vein). In initial descriptions, ILAs included ground-glass or reticular abnormalities, diffuse centrilobular nodularity, traction bronchiectasis, honeycombing, and non-emphysematous cysts (figure 1). In the definition proposed in this Position Paper, centrilobular nodularity, which is the typical presentation of smoking-related respiratory bronchiolitis,\(^2\) is not included as this feature is common, typically non-progressive, and not associated with fibrosis (panel 1).\(^2\) Although the 5% threshold is arbitrary and imprecise, it is retained to exclude minimal opacities and to conform to previous published literature. Focal or unilateral patchy ground-glass opacity seldom represents an ILD, and is classified as equivocal. Dependent abnormalities are regarded as equivocal unless persistent in the prone position. Pleuropulmonary fibroelastosis, sometimes an incidental finding on CT, is a clearly defined entity,\(^10\) which has not been included within ILAs in published series. Other findings not considered as ILAs are shown in panel 1 and figure 2.

Ensuring that specific descriptors of CT findings are provided in radiology reports is essential, as different imaging findings have very different implications. Relevant descriptors include craniocaudal and axial distribution and individual features, such as ground-glass or reticular abnormalities, traction bronchiectasis, architectural distortion, honeycombing, and non-emphysematous cysts (panel 1). Among these findings, the following subcategories are of prognostic significance: first, ground-glass opacity and reticular opacities without a predominant subpleural localisation; second, ground-glass opacity and reticular opacities with a predominant subpleural localisation without evidence of fibrosis; and finally, traction bronchiectasis, architectural distortion, and honeycombing, providing evidence of lung fibrosis.\(^2\) Non-emphysematous cysts, defined as luencies with irregular, well-defined walls, are often seen in cigarette smokers\(^3\) with or without other features of ILAs. These cysts can be distinguished from emphysema by the presence of a well-defined wall and from honeycombing by their irregular shape, varying size, and the absence of subpleural predominance.\(^4\) On histological analysis, non-emphysematous cysts usually correlate with airspace enlargement and fibrosis or smoking-related interstitial fibrosis, and might have prognostic significance, although they are not usually associated with imaging evidence of fibrosis.

ILAs with a non-subpleural distribution are usually non-progressive\(^5\) and not associated with increased mortality. Subpleural ILAs have potentially greater clinical significance and are further subcategorised according to the presence or absence of fibrosis (figure 1, panel 1). Fibrotic ILAs are associated with a higher rate of progression and death on 5-year follow-up.\(^6\) If fibrosis is present, the pattern can be further classified according to

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**Table: Interstitial lung abnormalities across study populations**

<table>
<thead>
<tr>
<th>Study characteristics</th>
<th>Population-based cohorts</th>
<th>Smoking and lung cancer screening cohorts</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MESA(^1)(^2)(^3)(^4)</td>
<td>Nagano, Japan(^5)(^6)</td>
</tr>
<tr>
<td>Total number of chest CT scans evaluated</td>
<td>3137</td>
<td>3061</td>
</tr>
<tr>
<td>Prevalence of ILAs</td>
<td>310 (10%)</td>
<td>80 (3%)</td>
</tr>
<tr>
<td>Mean age of those with ILAs (years)</td>
<td>75</td>
<td>62</td>
</tr>
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</table>

**Radiological progression**

<table>
<thead>
<tr>
<th>Overall progression, follow-up time</th>
<th>NA</th>
<th>46%</th>
<th>43%</th>
<th>63%</th>
<th>NA</th>
<th>20%</th>
<th>NA</th>
<th>20%</th>
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</tr>
</thead>
<tbody>
<tr>
<td>(hazard ratio [95% CI])</td>
<td>NA</td>
<td>(1·1–1·1)</td>
<td>(1·2–1·4)</td>
<td>(1·4–1·7)</td>
<td>NA</td>
<td>(1·2–2·0)</td>
<td>NA</td>
<td>(1·2–2·0)</td>
<td>NA</td>
</tr>
</tbody>
</table>

**Mortality**

<table>
<thead>
<tr>
<th>Relative risk of death, (hazard ratio [95% CI])</th>
<th>NA</th>
<th>NA</th>
<th>2·7</th>
<th>1·3</th>
<th>1·4</th>
<th>1·8</th>
<th>NA</th>
<th>2·0</th>
</tr>
</thead>
<tbody>
<tr>
<td>(hazard ratio [95% CI])</td>
<td>NA</td>
<td>(1·1–6·5)</td>
<td>(1·2–4·4)</td>
<td>(1·1–2·0)</td>
<td>(1·1–2·8)</td>
<td>(1·4–2·7)</td>
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</table>
Figure 1: Subcategories of interstitial lung abnormalities
(A) Non-subpleural and non-fibrotic. CT shows widespread ground-glass abnormality with central predominance (circled). (B) Subpleural non-fibrotic. CT shows predominantly subpleural ground-glass and linear abnormality without evidence of fibrosis (arrows). (C) Subpleural fibrotic. Traction bronchiectasis and architectural distortion are indicated by the ovals in the lingula and left lower lobe. This pattern would correspond to a probable usual interstitial pneumonia pattern.35,36

Panel 1: Definitions and subcategories of interstitial lung abnormalities
What are interstitial lung abnormalities (ILAs)?
- Incidental identification of non-dependent abnormalities, including ground-glass or reticular abnormalities, lung distortion, traction bronchiectasis, honeycombing, and non-emphysematous cysts
- Involving at least 5% of a lung zone (upper, middle, and lower lung zones are demarcated by the levels of the inferior aortic arch and right inferior pulmonary vein)
- In individuals in whom interstitial lung disease is not suspected

What are not ILAs?
- Imaging findings restricted to:
  - Dependent lung atelectasis
  - Focal paraspinal fibrosis in close contact with thoracic spine osteophytes (figure 2A)
  - Smoking-related centrilobular nodularity in the absence of other findings (figure 2B)
  - Mild focal or unilateral abnormality (figure 2C)
  - Interstitial oedema (eg, in heart failure)
  - Findings of aspiration (patchy ground-glass, tree in bud; figure 2C)

Preclinical and clinical identification:
- Preclinical interstitial abnormalities identified during screening of high-risk individuals (eg, those with rheumatoid arthritis, scleroderma, occupational exposure, familial interstitial lung disease)
- Findings in patients with known clinical interstitial lung disease

Subcategories of ILAs
- Non-subpleural: ILAs without predominant subpleural localisation (figure 1A)
- Subpleural non-fibrotic: ILAs with a predominant subpleural localisation and without evidence of fibrosis* (figure 1B)
- Subpleural fibrotic: ILAs with a predominant subpleural localisation and with evidence of pulmonary fibrosis* (figure 1C)

* Fibrosis is characterised by the presence of architectural distortion with traction bronchiectasis or honeycombing (or both).

the 2018 Fleischner and American Thoracic Society, European Respiratory Society, Japanese Respiratory Society, and Latin American Thoracic Society criteria as typical, probable, or indeterminate for usual interstitial pneumonia.37,38,39 About 2% of patients in the AGES-Reykjavík cohort had a probable or definite usual interstitial pneumonia pattern, were more likely to have subpleural ILA progression, and had worse survival compared with individuals without these patterns.39 It seems likely that fibrotic ILAs might be an important precursor to idiopathic pulmonary fibrosis (IPF) or other progressive fibrotic ILDs.

What CT protocol should be used to evaluate and follow up patients with ILAs?
When ILAs are detected, a dedicated chest CT examination could help to confirm and characterise the abnormality, especially if dependent atelectasis was present, if the initial scan of the lungs was incomplete (eg, an abdominal CT), or if the scan was done without thin sections, with an ultralow dose technique, or using first-generation, hybrid-type, iterative reconstruction methods, all of which might obscure fine lung details. On the dedicated CT examination (if indicated), thin sections (<1·5 mm) with moderate edge-enhancing reconstruction are helpful. Prone views are particularly important to distinguish dependent atelectasis and true interstitial abnormality, whereas expiratory imaging could potentially identify lobular air trapping as a clue to hypersensitivity pneumonitis. Potential recommended imaging protocols are outlined in 2018 guidelines.40 Subsequent Cts to evaluate for progression should use similar scanning protocols.

Pathological correlation of ILA
ILA is a radiological term with few published studies on pathological correlates. Pulmonary resection specimens for lung cancer in current or former cigarette smokers have a high frequency of background interstitial fibrosis. Katzenstein and colleagues41 reported a 60% prevalence of clinically occult fibrosis occupying more than 25% of the resected lobe. Most of these cases were viewed as smoking-related interstitial fibrosis, but usual interstitial pneumonia, pulmonary Langerhans’ cell histiocytosis, and asbestosis were also found. Similarly, in a larger study by Kawabata and colleagues,42 most cases of fibrosis were defined as airspace enlargement and fibrosis or respiratory bronchiolitis, the remainder being usual interstitial pneumonia. ILAs were not specifically identified on CT in these studies, but on the basis of the association between smoking and interstitial fibrosis it seems likely that many ILAs in smokers represent subclinical smoking-related fibrosis or macrophage accumulation.43–45 In 2018, Miller and colleagues46 evaluated histological correlates of ILAs in 424 lung nodule resections. Of 26 patients with ILAs, histology showed fibrosis in 19 (73%), with usual interstitial pneumonia in two (8%) individuals. Of note, 207 (52%) of 398 patients with no ILAs or an indeterminate status also showed histological fibrosis, suggesting that fibrosis can be below the resolution of imaging. Apart from usual interstitial pneumonia, the histological fibrosis seemed predominantly smoking related. In a similar study, Hung and colleagues47 found fibrotic ILD in 10% of 406 specimens from 397 patients, consisting of smoking-related interstitial fibrosis in 7%, usual interstitial pneumonia in 1%, non-specific interstitial pneumonia in 1%, and undefined in 1%. ILAs were present in 10% of cases with smoking-related interstitial fibrosis and in
100% with usual interstitial pneumonia. Similar to Miller and colleagues, Hung and colleagues found fibrotic changes in 51% of specimens with no radiological ILAs, although there were no cases of usual interstitial pneumonia in this category. A small number of cases with ILAs had granulomatous disease, non-specific interstitial pneumonia, undefined fibrosis, aspiration, or pulmonary Langerhans’ cell histiocytosis. Overall, these studies suggest that although usual interstitial pneumonia is sometimes present in patients with ILAs, a larger proportion of ILAs represent smoking-related changes. However, there is potential bias because the few studies published focus on findings in smokers or consist of lung cancer resections. Further study is needed to determine the frequency with which incidental histological fibrotic changes progress to clinically significant disease. To facilitate this research, reporting guidelines recommend that pathologists should record and categorise the presence in resection specimens of non-neoplastic lung parenchymal changes, such as emphysema, respiratory bronchiolitis, and interstitial fibrosis (with identification of a discernible pattern if possible).

**What are the risk factors for ILAs?**

Advanced age, a common feature of patients with IPF, is strongly associated with ILAs in almost all studies in which it has been assessed. For example, in smokers with and without COPD, each 10-year increase in age was associated with about a 2.2 times increase in the odds of detecting ILAs. Male sex has also been identified as a demographic risk factor in some studies of IPF and has been associated with ILAs in some, but not in all, cohorts. For example, in smokers with and without COPD, each male patient had about a 1.7 times increase in the odds of having ILAs compared with female patients.

Tobacco smoke exposure, commonly cited as an environmental risk factor for IPF as well as for other forms of ILD, is associated with ILAs in nearly all populations in which it has been evaluated. ILAs are associated with both the activity (e.g., current smoking status) and the quantity (e.g., pack years) of tobacco smoke exposure. For example, current smokers had about a 1.8 times increase in the odds of having ILAs compared with former smokers.

In a general population-based cohort, analyses of participants in MESA showed that self-reported occupational exposures to vapours, gases, dusts, and fumes were associated with an increased prevalence of high-attenuation areas and ILAs. In the MESA-Lung study, increased exposure to nitrogen oxides, a marker of exposure to traffic-related air pollution, was also associated with ILAs. Similarly, in the Framingham Heart Study, elemental carbon exposure (another common metric of traffic-related air pollution) was associated with ILAs and ILA progression.

The most consistent genetic risk for both IPF and familial interstitial pneumonia has been increased copies of the minor allele of a common variant in the promoter of the MUC5B gene (rs35705950). Similarly, the associations between the MUC5B promoter genotype, ILAs, and ILA progression have been consistently replicated. For example, in the Framingham Heart Study, COPDGene study, and AGES-Reykjavik cohorts, each copy of the minor allele of the MUC5B promoter polymorphism is associated with between a 1.5 and 2.7 times increase in the risk of presenting with ILAs, particularly in those with subpleural ILA (e.g, the MUC5B promoter genotype is more strongly associated with subpleural abnormalities than with centrilobular nodules). Although additional overlapping findings of genetic association have been shown in comparisons of genome-wide association analyses of IPF and ILAs, novel genetic association with ILAs suggests that disorders other than IPF are also likely to be present among some research participants with these imaging abnormalities.

**What are the clinical outcomes of ILAs?**

ILAs have been associated with adverse clinical outcomes in numerous populations. These include general population cohorts and populations of smokers enriched for the presence of COPD or undergoing lung cancer screening.

**Progression of ILAs**

Estimates of the rate of imaging progression of ILAs range from 20% over 2 years in the National Lung Screening Trial to 48% over 5 years in the AGES-Reykjavik study (figure 3). Thus, although not all cases of ILAs progress, progression is more likely to be detected when followed up over longer time periods. Additionally, patients with ILAs without clear pulmonary fibrosis might subsequently develop traction bronchiectasis, honeycombing, or patterns consistent with usual interstitial pneumonia. However, the proportion of such cases that evolve to usual interstitial pneumonia on long-term follow-up remains unclear.
Specific imaging features and patterns can identify ILAs that are most likely to progress over a 5-year interval. For example, in a study by Putman and colleagues, patients with subpleural reticular changes, lower lobe predominant changes, or traction bronchiectasis had more than six times increase in their odds of imaging progression than those with ILAs without these features, even after adjusting for important covariates (e.g., age and smoking history). In that study, all cases of honeycombing progressed over 5 years. Conversely, the presence of centriflobular nodules was associated with ILAs that were unlikely to progress.

ILA progression and lung function decline were explored in a Framingham Heart Study cohort. Progression (including both the development of new ILAs and the progression of existing ILAs) occurred in 6% of the population over approximately 6 years. Patients with imaging progression in the Framingham Heart Study had an accelerated decline in forced vital capacity (FVC) compared with those patients without ILAs or those with ILAs that did not progress. However, the annual decline in FVC in patients with ILA progression in the Framingham Heart Study (about 60 mL per year) with an excess annual decline of about 30 mL per year compared with those without ILAs) was substantially less than the annual decline in FVC generally noted among patients with IPF (approximately 200 mL per year). Whether the excess FVC decline associated with ILA progression on imaging represents a small subgroup with major FVC decline (averaging to a small FVC decline across all progressing patients) or a larger subgroup with subclinical disease that tends to be less pronounced than clinically apparent IPF is not clear.

**Mortality**

One of the most consistent findings with regard to ILAs is the association with increased mortality, both in general population samples and among populations of smokers enriched for COPD or undergoing lung cancer screening (table). In the Framingham Heart Study and AGES-Reykjavik cohorts, this increase in mortality was most strongly associated with imaging progression of ILAs. In the AGES-Reykjavik cohort, specific imaging patterns indicative of pulmonary fibrosis were associated with earlier mortality. In addition to increased all-cause mortality, ILAs were associated with increased respiratory mortality in the AGES-Reykjavik cohort. From a Brigham and Women’s Hospital cohort of patients with systemic inflammatory response syndrome or sepsis, ILAs were associated with increased rates of acute respiratory distress syndrome and increased inhospital mortality. ILAs are also associated with increased mortality in patients with COPD and lung cancer, and in individuals who undergo transcatheter aortic valve replacement. Increased quantitative metrics of ILAs (based on an increased number of high-attenuation areas of the lung and local histogram-based methods designed to identify ILAs) are also associated with increased mortality.

Although ILAs have been associated with increased mortality from pulmonary fibrosis, it is important to recognise that the contribution of ILAs to these elevated mortality rates far exceeds the expected rate of progression to clinically detectable ILD. It is also important to note that respiratory-related deaths, which were more common among those with ILAs in the AGES-Reykjavik cohort, were reported in less than 15% of those with ILAs. This observation suggests that although some of the association between ILAs and death could be due to pulmonary fibrosis, those patients with ILAs could possibly be at an excess risk of death because of accelerated physiological ageing or other causes of death that are not directly related to pulmonary disease.

**Lung cancer mortality and treatment toxicity**

Several studies have shown an association between pretreatment ILAs and cancer-associated mortality, including patients with early stage cancer treated with surgical resection, as well as patients with advanced stage 4 disease treated with systemic therapy. The cause of increased mortality is not clear, but other studies suggest that lung injury risk associated with ILAs and cancer therapies might be important. Specifically, lung irradiation and systemic treatment with chemotherapy and targeted tyrosine kinase inhibitors, immunotherapy checkpoint inhibitors, and antibody–drug conjugates are associated with an increased risk of pneumonitis in patients with pre-existing ILAs (figure 4). Pre-existing ILAs increase the risk of extensive radiation pneumonitis in patients with early stage lung cancer treated with stereotactic body radiotherapy and in patients with small-cell lung cancer treated with 50–60 Gy of thoracic radiotherapy.

Immune checkpoint inhibitors have emerged as standard first-line therapy for patients with advanced non-small-cell lung cancer and for other malignancies. Overall, the rate of immunotherapy-associated pneumonitis is approximately 5%, and this toxicity is manageable when
recognised early and treated appropriately in accordance with the National Cancer Institute’s Common Terminology Criteria for Adverse Events grade at presentation.\textsuperscript{7,8} Evidence indicates that pneumonitis risk is increased by ILAs. Nakanishi and colleagues\textsuperscript{9} examined pretreatment chest CT scans for ILAs in 83 patients treated with the anti PD-1 antibodies nivolumab or pembrolizumab. The incidence of immunotherapy-associated ILD was high at 17% (n=14). Multivariate analysis showed that pre-existing ILAs were associated with a six times increase in the risk of drug-associated ILD, with a predominant ground-glass pattern of pneumonitis. Given the life-threatening nature of malignancies treated with immune checkpoint inhibitors, the benefits of the therapy, and the undefined risks associated with ILAs, clinicians should discuss the possible increased risk of pneumonitis with patients who have ILAs. Additionally, clinicians should consider active monitoring for symptoms, physiological alterations, and radiological progression of drug-associated pneumonitis (figure 5).

How should ILAs be evaluated and monitored?

To date, only minimal evidence exists to support a specific management plan for ILAs. The following proposal is based on the available published literature and the consensus clinical opinion of the authors. The first goal is to separate those patients with current clinically significant disease from individuals who might be at risk of developing disease. This distinction could be established by a series of questions that incorporate a general approach to ILD (figure 5). In all patients, a standard evaluation of potential explanations for the presence of ILAs should occur, including factors such as cigarette smoking or other inhaled exposures, drug toxicity, systemic disease (eg, occult connective tissue disorders), or recurrent aspiration of oroesophageal contents. Individuals with respiratory symptoms or signs, clinically relevant pulmonary physiological or gas transfer abnormalities, or extensive CT abnormality (disease involving three or more of the six lung zones consisting of the right and left upper, middle, and lower lung zones) should be referred for pulmonary evaluation, ideally with access to multidisciplinary discussion. Management of patients with a diagnosed ILD should follow standard guidelines.

Once ILD is excluded, ILAs can be separated into those at higher risk of progression to ILD and those at lower risk. Risk factors for progression include cigarette smoking, other inhalational exposures, medications, physiological or gas exchange findings not felt to reach the threshold of clinical significance, and specific radiological features such as evidence of fibrosis and subpleural, basal predominant distribution (panel 2).

Follow-up of patients with ILAs can be based on the presence of risk factors for progression. Individuals without risk factors should be advised to return for evaluation if they develop symptoms of respiratory impairment. For example, non-subpleural ILAs seldom progress, and individuals with only these findings can be followed up expectantly. In individuals with one or more risk factors, systematic follow-up should be considered. The appropriate timing of repeated clinical evaluation (including a focused history and chest exam, chest
imaging, pulmonary physiology, and gas exchange) is unknown. In the absence of prospective data, clinical experience suggests that a first follow-up at 3–12 months to look for symptomatic or physiological progression is probably appropriate in most patients at increased risk. Individuals with ILAs are likely to benefit from additional clinical follow-up beyond 1 year, but the optimal frequency and duration of follow-up is unknown. Similarly, the optimal interval for follow-up CT scans is unknown, but might include a follow-up scan at 12–24 months or sooner in patients who develop symptoms or impaired pulmonary function. Progression can be defined by the development of clinically significant respiratory symptoms and signs (eg, the new presence of exercise limitation or characteristic crackles on auscultation, or both), the development of abnormal pulmonary physiology or gas exchange (or a clinically significant decline in normal values), or an increase in the extent of CT abnormalities, particularly with the development of specific fibrotic features. Optimal management of progressive ILAs is unknown, so this patient group might be an appropriate population for a prospective treatment trial.

In patients with ILAs undergoing surgery or other therapy, caution should be exercised because they appear to be at increased risk of rapid disease acceleration or an acute exacerbation. The clinician should regard ILAs as an important comorbidity that should be considered in planning treatment and subsequent monitoring. Because positive pressure ventilation might be a risk factor for developing acute respiratory distress in patients with ILAs, a low-volume, low-pressure ventilatory approach should be considered for those needing mechanical ventilation. Medications that are known to cause ILD should be avoided if possible.

**What is the role of quantitative evaluation?**

Methods for quantitative evaluation of ILAs include assessment of the proportion of high-attenuation areas, local histogram evaluation, and deep learning-based textural evaluation. Automatic quantification of CT density of the lungs has been used to identify the proportion of lung voxels with high-attenuation areas, typically between –600 and –250 Hounsfield units (the normal CT attenuation of the lung is about –750 Hounsfield units).12,21 High-attenuation areas are associated with elevated serum concentrations of inflammatory biomarkers, reduced FVC and exercise capacity, and higher mortality (including higher mortality from ILD).12,21 The presence of high-attenuation areas is also associated with a higher prevalence of ILAs at follow-up CT.21 Despite these clear epidemiological associations, high-attenuation areas appear to be neither sensitive nor specific for subsequent appearance of ILAs.12 For this reason, the use of the term subclinical ILD as a synonym for increased high-attenuation areas is not recommended. The clinical significance of high-attenuation areas remains unclear, and assessment in individual patients is limited by the multiple technical and patient-related reasons for an elevated proportion of high-attenuation areas, including scanner variation, inadequate inspiration, obesity, and pulmonary atelectasis.12

Quantitative imaging provides an objective recognition of regional disease pattern of the lung that can increase the diagnostic reliability and severity assessment of ILD. Computer-based CT approaches to identify interstitial

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**Panel 2: Risk factors for progression of interstitial lung abnormalities**

**Clinical risk factors**

- Cigarette smoking
- Other inhalational exposures
- Medications (eg, chemotherapy, immune checkpoint inhibitors)
- Radiation therapy
- Thoracic surgery
- Physiological or gas exchange findings at lower limits of normal

**Radiological risk factors**

- Non-fibrotic interstitial lung abnormalities (ILAs) with basal and peripheral predominance
- Fibrotic ILAs with basal and peripheral predominance but without honeycombing (ILAs with probable usual interstitial pneumonia pattern)
- Fibrotic ILAs with basal and peripheral predominance and honeycombing (ILAs with usual interstitial pneumonia pattern)

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**Figure 6:** Computer-based classification of interstitial lung abnormalities with the histogram approach

(A) CT images with subpleural non-fibrotic interstitial lung abnormalities and emphysema in a participant in the COPDGene study. (B) CT image overlays of computer-based classification of interstitial lung abnormalities using artificial intelligence, showing objective quantification of different injury patterns. Regions of interstitial lung abnormality are shown in blue. Normal parenchyma (red), emphysema (green), and paraseptal emphysema (yellow) are also subtyped.16
subtypes are based on density histogram analysis, texture-based analysis, and deep learning approaches.\(^{30-37}\) In general, these approaches are sensitive enough to detect ILD in patients at high risk and provide a more reproducible assessment than visual CT scoring.\(^7\) For ILAs, individuals with a lower percentage of normal lung by local histogram measurements had a higher prevalence of clinical impairment, poorer quality of life, higher risk of death, and association with the common variant in the promoter of the MUC5B gene (figure 6).\(^{17}\) Using a different local histogram-based system in 217 individuals undergoing resection for lung cancer, the fibrosis score correlated with the presence of ILAs and was an independent predictor of decreased disease-free survival.\(^{3}\) A study in family members of individuals with familial pulmonary fibrosis showed that data-driven texture analysis could detect early interstitial changes with 84% sensitivity and 86% specificity (figure 7).\(^{49}\) However, the role of quantitative CT as a screening tool for ILAs requires further validation. Potential sources of variation in quantitative imaging of ILAs include dependence on training data, variation in inspired lung volumes, sensitivity to image noise from CT acquisition dose, vendor differences and reconstruction method, and variation in segmenting the subpleural fibrotic lung from the chest wall. Annotated datasets are needed to provide a reference benchmark to establish the robustness of each approach. Considering that the characteristics of ILAs are subtle and varied, the stability of assessment by computer-based analysis should be tested, improved, and applied in further studies. New advances in artificial intelligence and deep learning might overcome some of these limitations.\(^{30,39-44}\)

**Outline of future research needs and priorities**

The preliminary radiological criteria for ILAs presented in this Position Paper require robust evaluation to determine their reproducibility and application to clinical practice, including in lung cancer screening (panel 3). The effect of the proposed management plan on intermediate and long-term outcomes must be evaluated (figure 5). To understand the prevalence and natural course of ILAs in the lung cancer screening population, ILAs should be considered as a specific subcategory under the significant other findings modifier in the LungRADS scoring system,\(^7\) as used in the USA; the Korean Society of Radiology has already implemented a similar change.\(^{68,69}\) In addition to clarifying criteria for visual evaluation of ILAs, quantitative CT methods for evaluation of disease extent and determination of progression will need to be developed and validated. An important element will be the determination of optimal thresholds on visual and quantitative CT that define significant disease by predicting significant physiological progression, the development of clinically significant disease, and mortality. Since antifibrotic treatment slows the rate of physiological progression in patients with IPF,\(^{49,50}\) and in other forms of progressive lung fibrosis,\(^{50,51}\) it is possible that early treatment in a high-risk population with ILAs might reduce the rate of progression. However, there is a clear need for further epidemiological, biomarker, and machine learning studies in existing and novel cohorts to identify groups at higher risk of progression and to understand the trajectory of progression of ILAs to clinically significant pulmonary fibrosis. The availability of this information would potentially support the design of future clinical trials in higher-risk individuals.

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**Panel 3: Key uncertainties with interstitial lung abnormalities**

- Reproducibility of radiological criteria
- Validity and efficacy of follow-up regimen
- Prevalence in the lung cancer screening population
- Prevalence in younger cohorts
- Risk factors for progression
- Natural history of non-fibrotic interstitial lung abnormalities (ILAs)
- Optimal extent thresholds for predicting significant physiological progression, development of clinically significant disease, and mortality by visual and quantitative CT evaluation
- Importance of incidentally identified histological evidence of interstitial abnormality
- Quantitative techniques: predictive value for adverse outcome, technical variability, and inter-patient variability
- Role of biomarkers in predicting progression
- Strategy for cohort enrichment in clinical trials for patients with ILAs that are likely to progress
- Risk factors and preventive strategies for complications of cancer treatment surgery, chemotherapy, immunotherapy, and radiotherapy in patients with ILAs

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**Figure 7: Quantification of progression for interstitial lung abnormalities with data-driven texture analysis**

(A) A baseline CT scan shows subpleural non-fibrotic interstitial lung abnormalities with fibrotic changes. (B) CT 5 years later shows clear progression. (C) Baseline data-driven textural analysis shows overall extent of fibrosis as 1.5% (red). (D) Data-driven textural analysis of follow-up scan at 5 years shows that the extent of fibrosis increased to 4.6% (red).

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**Key uncertainties with interstitial lung abnormalities**

- Reproducibility of radiological criteria
- Validity and efficacy of follow-up regimen
- Prevalence in the lung cancer screening population
- Prevalence in younger cohorts
- Risk factors for progression
- Natural history of non-fibrotic interstitial lung abnormalities (ILAs)
- Optimal extent thresholds for predicting significant physiological progression, development of clinically significant disease, and mortality by visual and quantitative CT evaluation
- Importance of incidentally identified histological evidence of interstitial abnormality
- Quantitative techniques: predictive value for adverse outcome, technical variability, and inter-patient variability
- Role of biomarkers in predicting progression
- Strategy for cohort enrichment in clinical trials for patients with ILAs that are likely to progress
- Risk factors and preventive strategies for complications of cancer treatment surgery, chemotherapy, immunotherapy, and radiotherapy in patients with ILAs

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The importance of ILAs as a risk factor for complications in the treatment of lung cancer requires further evaluation, for example, in clinical trials of checkpoint inhibitors with a focus on prevention and management of acute pneumonitis. This analysis could be done retrospectively from existing CT datasets in clinical trials and could inform a prospective comparison of specific approaches to preventing progression in the context of thoracic surgery, radiotherapy, and chemotherapy. The role of ILAs as a predictor of acute interstitial pneumonia also merits further evaluation, given that unsuspected usual interstitial pneumonia has been identified in 50% of patients who die with acute interstitial pneumonia.\(^6\) We have excluded preclinical interstitial abnormalities identified during the screening of individuals at high risk (eg, those with rheumatoid arthritis, scleroderma, occupational exposure, and familial interstitial lung disease) from the scope of this Position Paper to simplify the approaches. However, future studies and discussions are needed to investigate the role of ILAs in preclinical interstitial abnormalities identified during screening of individuals at high risk.\(^*\)

Further study is needed to determine the importance of incidentally found histological fibrosis to determine which cases are more likely to progress to clinically significant disease. In particular, there is an opportunity to clarify the effect of specific histological findings and cellular subpopulations on long-term outcome. Additional related questions include the natural history and biological cause of non-fibrotic ILAs, of smoking-related interstitial fibrosis, and of pleuroparenchymal fibroelastosis. There is also a need for better understanding of risk factors for the development of ILAs. Finally, the recognition of ILAs offers exciting opportunities for identifying pathogenetic abnormalities in early pulmonary fibrosis and early usual interstitial pneumonia; sequential evaluation of biomarkers in individuals with ILAs might help to identify biological abnormalities that predispose to subsequent development of IPF.

**Conclusions**

ILAs are important because they are associated with mortality as well as increased risk of complications from surgery, chemotherapy, and radiotherapy. Separating clinically significant ILD from ILAs is essential. The morphology and distribution of ILAs should be clearly described and the descriptive categories of non-subpleural, subpleural non-fibrotic, and subpleural fibrotic ILAs should be recorded in the radiology report, as this information could be useful in predicting progression and mortality (panel 4). Risk factors for ILAs include age, cigarette smoking and other inhalational exposures, and genetic markers. Although this Position Paper proposes a rational strategy that can help to identify when ILAs are likely to represent clinically significant ILD, future work is needed to determine the best approach to follow up ILAs in individuals in whom the evaluation is less definitive. We believe that establishing a common terminology for communication and a clear understanding of current knowledge are important steps towards further advances in the multidisciplinary approach to ILAs.

**Contributors**

DAL developed and implemented the systematic search strategy. CJR advised on the systematic search. All authors participated in the

**Search strategy and selection criteria**

A medical librarian searched in Medline, Embase, Cochrane Central Registry of Controlled Trials, and the Health Technology Assessment database to identify publications related to interstitial lung abnormalities. We included studies from database inception through to Feb 13, 2019, and restricted to English language. Details of the search strategy are provided in the appendix (pp 1–3). Key search terms were “interstitial”, “lung”, “abnormal”, and “subclinical or pre-clinical”. The literature search resulted in 700 references, of which 616 were excluded (duplicates \(n=11\) and 605 references with little relevance to the key questions based on screening of the reference title \(n=455\) or the reference abstract \(n=150\)), yielding 84 manuscripts that underwent review for inclusion. Review of the text found that 60 of these manuscripts were not relevant to the key questions, resulting in 24 references that were analysed for the final Position Paper. Additional references were added by members of the writing group.

**Panel 4: Recommendations for the evaluation and reporting of interstitial lung abnormalities**

**CT protocol**

- Thin sections (<1·5 mm) are essential
- Prone and expiratory scans might be necessary to confirm and characterise interstitial lung abnormalities (ILAs)

**CT description**

- Axial and craniocaudal distribution
- CT findings: including ground-glass abnormality, reticular abnormality, traction bronchiectasis, honeycombing, and cysts
- CT category: non-subpleural ILA, subpleural non-fibrotic ILA, or subpleural fibrotic ILA

**Clinical evaluation**

- Distinguish ILAs from clinically significant interstitial lung disease (figure 5)
- Identify risk factors for progression (panel 2)
- Follow-up evaluation (figure 5)

**Pathology evaluation**

- On lung cancer resections, assess background lung from cancer resections and document histological patterns diagnostic of suspicion for interstitial lung disease
- Review such cases in a multidisciplinary team setting to determine whether ILAs or clinically significant interstitial lung disease is present
literature search. HH, GMH, LR, KKB, AUW, MR-J, AGN, MBB, DCC, RSKJ, CAP, KSI, YI, and DAL wrote the first draft of the Position Paper. All authors critically reviewed the manuscript and approved the final version, taking accountability for the work. The Document Development and Oversight Committee and the Executive Committee of the Fleischner Society approved the manuscript before submission to the journal.

Declaration of interests
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