Evaluation of patients with fibrotic interstitial lung disease: A Canadian Thoracic Society position statement

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ABSTRACT
The evaluation of a patient with fibrotic interstitial lung disease (ILD) includes assessment of clinical, radiological, and often histopathological data. There are currently no specific recommendations to guide the evaluation of a patient with fibrotic ILD within the context of the Canadian practice landscape. This position statement from a multidisciplinary panel of ILD experts provides guidance related to the diagnostic modalities commonly used in the evaluation of fibrotic ILD, including radiological studies, histopathologic sampling, assessment for rheumatologic disease and need for evaluation in a multidisciplinary setting. Key messages are provided to guide clinical practice based on a thorough review of the scientific literature.

RESUMÉ
L’évaluation d’un patient atteint de fibrose pulmonaire interstitielle comprend l’évaluation de données cliniques, radiologiques et souvent histopathologiques. Il n’existe actuellement aucune recommandation pour guider l’évaluation d’un patient atteint de fibrose pulmonaire interstitielle dans le cadre de la pratique canadienne. Cet énoncé de position d’un comité multidisciplinaire d’experts en maladie pulmonaire interstitielle donne des conseils relativement aux modalités de diagnostic communément utilisées dans l’évaluation de la fibrose pulmonaire interstitielle, y compris les études radiologiques, l’échantillonnage histopathologique et l’évaluation pour un trouble rhumatologique, de même que la nécessité que l’évaluation soit menée dans une perspective multidisciplinaire. Afin de guider la pratique clinique, des messages clés fondés sur une revue approfondie de la littérature sont énoncés.

Introduction
The interstitial lung diseases (ILDs) are a heterogeneous group of disorders characterized by inflammation and/or fibrosis of the pulmonary interstitium. ILDs are challenging to diagnose, with over 200 unique subtypes that frequently have overlapping clinical and radiological features. The fibrotic ILDs include those with radiological and/or pathological evidence of fibrosis and are typically characterized by progressive physiologic impairment, dyspnea, functional limitation and early mortality.

Current diagnostic guidelines are based on limited evidence and focus on idiopathic pulmonary fibrosis (IPF) or other idiopathic interstitial pneumonias (IIPs); however, a more general approach is needed at the time of initial assessment. A recent survey of Canadian pulmonologists revealed heterogeneous practice patterns and deviations from international consensus recommendations. These issues indicate the need for recommendations applicable to a broader variety of fibrotic ILDs.

The objectives of this position paper are to:
1) Summarize the available literature on topics relevant to the evaluation of patients with fibrotic ILD.
2) Provide evidence- and expertise-based key messages for the evaluation of patients with fibrotic ILD.

There are several essential components to the evaluation of fibrotic ILD that should be addressed in all patients. These include a thorough history identifying symptoms suggestive of connective tissue disease and relevant occupational and/or environmental exposures, a family history of ILD or autoimmune disease, a detailed physical examination and pulmonary function testing. These and other “good clinical practice” items...
were not reviewed as these should be evaluated in all patients with fibrotic ILD, and a comprehensive review of each of these topics is beyond the scope of the current work. In contrast, there are additional items that have insufficient data to support evidence-based key messages (eg, specific circulating antibodies in the evaluation of hypersensitivity pneumonitis, genetic testing in the routine evaluation of a patient with fibrotic ILD), and these topics will similarly not be reviewed.

The key messages provided herein are intended to guide evidence-based decision-making by clinical care providers (eg, general internists, pulmonologists, radiologists, pathologists, thoracic surgeons and any others initiating the evaluation for fibrotic ILD) and should be interpreted in the context of patient wishes and preferences. Other individuals, including family members and/or allied health care providers, should be involved in these discussions when appropriate to address the unique needs of each patient.

Methods
A working group was created within the Canadian Thoracic Society’s Clinical Assembly on Interstitial Lung Disease. The group was co-chaired by two authors (KAJ and CJR), and included 9 adult pulmonologists with expertise in ILD patient care, 1 rheumatologist, 2 chest radiologists, 2 lung pathologists and 1 thoracic surgeon.

The document was developed in accordance with Canadian Thoracic Society (CTS) requirements for a position statement. The working group identified 7 clinically relevant questions pertaining to the diagnosis of fibrotic ILD and then conducted a narrative review of the scientific literature on these topics. Subcommittees produced a literature summary and identified key messages for each question, agreed upon through collaborative discussion by the working group. The completed document was reviewed by 2 ILD experts external to the CTS, 1 member of other CTS guideline committees, and members of the CTS Canadian Respiratory Guidelines Committee reviewed the completed document. One member completed the AGREE II checklist. Original reviews and responses to reviews are posted along with the guideline and all author conflicts of interest, at respiratoryguidelines.ca. The CTS Executive approved the final document for publication. The position statement will be updated in accordance with the CTS Living Guideline Model (www.respiratoryguidelines.ca).

Summary of evidence and key messages

High resolution computed tomography

What are the appropriate specifications for computed tomography (CT) in the evaluation of fibrotic ILD?
High-resolution CT (HRCT) cross-sectional chest imaging permits visualization of the lungs in fine anatomic detail. HRCT is a critical component of the evaluation of the ILDs and should be performed in all patients with suspected fibrotic ILD, unless contra-indicated. The HRCT protocol should include thin-collimation axial scans or thin-section reconstruction of volumetric helically acquired data using multi-detector CT and narrow detector width (0.5-1.25 mm), image reconstruction with a high spatial frequency (sharp or edge-detecting) algorithm and sufficient radiation to keep image noise and patient exposure at acceptably low levels.

HRCT studies should be performed at full inspiration. Expiratory imaging should be considered, particularly when there is suspicion of diseases that can cause air trapping (eg, hypersensitivity pneumonitis, obliterative bronchiolitis). Prone imaging can help differentiate atelectasis from ILD in patients with mild subpleural changes in dependent parts of the lungs. Volumetric acquisition provides complete imaging of the lung, contiguous imaging of specific lesions (eg, nodules), multiplanar reformatting, precise comparison to prior exams and assessment of additional lung abnormalities. Contrast enhancement may be considered in cases where there is suspected concomitant thromboembolic disease, lymph node enlargement, mass lesions or pleural disease. Low dose chest CT for lung cancer screening or pulmonary nodule assessment can identify abnormalities suggestive of ILD but does not eliminate the need for further imaging with HRCT.

The diagnostic accuracy of HRCT depends on the experience of the interpreting radiologists and whether interpretation is provided within the context of a multidisciplinary discussion (MDD). The high inter-observer agreement among academic radiologists suggests the potential for greater diagnostic accuracy; however, the specific clinical implications of this finding are currently unknown. Academic radiologists have higher inter-observer agreement than community radiologists for the interpretation of HRCT for ILD, with the latter having a greater tendency to report a usual interstitial pneumonia (UIP) pattern. Diagnostic accuracy can be as high as 90% when an experienced chest radiologist confidently identified a UIP pattern, and this can eliminate the need for surgical lung biopsy in some patients.

Key Messages: High Resolution Computed Tomography
- All patients being evaluated for fibrotic ILD should undergo high-resolution computed tomography (HRCT) imaging of the chest, unless contraindicated.
- Expiratory and prone HRCT imaging may provide valuable diagnostic information.
- Continuous images with 0.5-1.25 mm slice thickness are the standard of care for the imaging evaluation of fibrotic ILD.
- Chest HRCT should be interpreted in the context of a multidisciplinary discussion in cases with diagnostic uncertainty.

Evaluation for connective tissue disease

Which autoimmune serologic tests should be performed in the evaluation of fibrotic ILD?
Connective tissue diseases (CTDs) are systemic autoimmune disorders that typically affect the skin and musculoskeletal system. ILD is an important cause of morbidity and mortality in patients with CTD, commonly occurring in patients with systemic sclerosis, rheumatoid arthritis, idiopathic inflammatory myopathies, mixed CTD, Sjögren’s syndrome and systemic lupus erythematosus. Consensus criteria have recently been proposed for interstitial pneumonia with autoimmune features (IPAF), defined as patients with ILD who have autoimmune features that do not meet criteria for a defined CTD.
Additional studies are required to determine whether it is appropriate to consider this population a distinct clinical entity.

It is important to identify CTD as an underlying cause of ILD given the different management and prognosis compared to other ILD subtypes. Most importantly, patients with CTD-ILD may benefit from immunosuppressive agents that carry an increased risk of death in patients with IPF. Detailed evaluation for symptoms and signs of underlying CTD should be performed at baseline in all patients with fibrotic ILD, and should be repeated during follow-up since ILD can also develop prior to overt features of CTD. No consensus exists on which autoimmune serology should be routinely performed when screening a broader ILD population for the presence of CTD; however, recent IPF guidelines recommend obtaining a rheumatoid factor (RF), antinuclear antibody (ANA) and anti-cyclic citrullinated peptide (CCP) antibody in patients with suspected IPF. It is likely appropriate to perform these studies in all fibrotic ILD patients, with additional targeted serologic testing reserved for patients with a high pre-test probability of CTD. Indirect immunofluorescence is more sensitive than solid phase assays for detecting ANA, and should be used for the initial screening test.

Key messages: Autoimmune Serology Testing

- All patients with fibrotic ILD should undergo baseline anti-nuclear antibody testing (by indirect immunofluorescence), rheumatoid factor, and anti-cyclic citrullinated peptide antibody testing, with additional serologic testing in patients with a high pre-test probability for connective tissue disease.
- Additional research is required to determine whether "interstitial pneumonia with autoimmune features" (patients with ILD who have autoimmune features that do not meet criteria for a defined connective tissue disease) is a distinct clinical entity.

When should a patient with fibrotic ILD be referred to a rheumatologist?

Circulating low-level autoantibodies have been reported in patients with IPF who lack clinical findings of CTD. An abnormal ANA is present in 25%-41% of IPF patients at a titre ≥1:40, and an elevated RF present in 6%-7%. However, the significance of these autoantibodies is unclear given the similar frequencies seen in healthy age-matched control populations and the similar prognostic of serology-positive and serology-negative patients with IPF. The mere presence of an autoantibody is therefore insufficient to provide a CTD-ILD diagnosis without additional confirmatory clinical data.

There are limited data to guide the need for assessment by a rheumatologist in a patient with fibrotic ILD. In the absence of additional CTD features, ILD patients with low-titre ANA and/or RF likely do not require referral to a rheumatologist, particularly if the onset of ILD is after the age of 65 years. Conversely, referral to a rheumatologist should be considered in patients with suggestive clinical features, specific autoantibodies, or high titre ANA (>1:320) and/or RF (>60 IU/mL). Similarly, a rheumatology referral should be considered in patients that meet criteria for IPAF (see Reference 19 for IPAF criteria). Some Canadian centres offer combined pulmonary-rheumatology ILD clinics for coordinated multidisciplinary evaluation of patients with suspected CTD-ILD.

Key messages: Bronchoalveolar Lavage

- Bronchoalveolar lavage can be diagnostic of some pulmonary diseases in the appropriate clinical context, including infection, malignancy and some nonfibrotic ILDs.
- It is unclear whether bronchoalveolar lavage adds incremental diagnostic information beyond a thorough evaluation for alternative diagnoses in patients with fibrotic ILD.

The uncertain benefit of BAL should be balanced against the potential risks, including transient worsening of hypoxemia and, rarely, acute exacerbation of the underlying ILD. Recent guidelines recommend against BAL in the evaluation of most IPF patients but acknowledge that it may be appropriate in a minority. Given the absence of additional supportive data, the primary role of BAL in fibrotic ILD is to exclude infection in patients presenting with suggestive clinical and radiological features, especially if immunosuppressive therapy is being considered.

Key messages: Rheumatology Referral

- A low-titre autoantibody may not be clinically significant and may not mandate rheumatology referral in the absence of other features of connective tissue disease.
- ILD patients with suggestive clinical features, specific autoantibodies, or high titre anti-nuclear antibody (>1:320) and/or rheumatoid factor (>60 IU/mL) should be referred to a rheumatologist.

Bronchoalveolar lavage

When should bronchoalveolar lavage be performed in the evaluation of fibrotic ILD?

Bronchoalveolar lavage (BAL) is a minimally invasive and generally well-tolerated method to collect cells, organisms and inhaled particles from the lower respiratory tract and alveolar spaces. The recommended techniques for obtaining and processing BAL fluid have been described elsewhere and are essential to ensure meaningful results. BAL can be diagnostic in many pulmonary diseases in the appropriate clinical context, including infection, malignancy and some nonfibrotic ILDs (eg, pulmonary alveolar proteinosis, eosinophilic pneumonia).

There are limited data supporting the diagnostic utility of BAL in fibrotic ILD. BAL neutrophilia >5% is most consistent with IPF although can be found in other fibrotic ILDs, whereas lymphocytosis >25% suggests granulomatous disease and >50% is highly suggestive of hypersensitivity pneumonitis (HP) or idiopathic nonspecific interstitial pneumonia (NSIP). BAL cellular analysis, and particularly a lymphocytosis >30%, resulted in a change in diagnosis for 8% of patients that had a confident diagnosis of IPF based on clinical and radiological features in a previous cohort study. Other studies have conversely shown that significant lymphocytosis is less common in chronic (fibrotic) HP compared to acute or subacute forms. Some reports suggest an informative role of lymphocyte subset analysis (eg, CD4 to CD8 ratio) in sarcoidosis, but this test has no clear utility in fibrotic ILD. There are no adequately designed studies evaluating whether BAL adds incremental diagnostic information beyond a thorough evaluation for alternative diagnoses in patients with fibrotic ILD.

The uncertain benefit of BAL should be balanced against the potential risks, including transient worsening of hypoxemia and, rarely, acute exacerbation of the underlying ILD. Recent guidelines recommend against BAL in the evaluation of most IPF patients but acknowledge that it may be appropriate in a minority. Given the absence of additional supportive data, the primary role of BAL in fibrotic ILD is to exclude infection in patients presenting with suggestive clinical and radiological features, especially if immunosuppressive therapy is being considered.
Transbronchial biopsy

When should transbronchial biopsy be performed in the evaluation of fibrotic ILD?

Transbronchial biopsy (TBBx) provides small (~1-3mm²) samples of parenchymal lung tissue taken through the working channel of a bronchoscope. TBBx may have a role in the evaluation of some nonfibrotic ILDs including sarcoidosis,11,12 however, its role in the evaluation of fibrotic ILD is less clear. In one study, 3/33 patients (9%) with bird fancier’s disease had the classic triad of HP findings on TBBx (lymphocytic-histiocytic infiltrate, poorly formed granuloma, bronchiolitis obliterans), although poorly formed granulomas were identified in 21% of cases.13 TBBx is less helpful in diagnosing other fibrotic ILDs, and can also provide misleading information. For example, TBBx showed an NSIP pattern in 9/18 patients (50%) with usual interstitial pneumonia (UIP) pattern proven by surgical lung biopsy.43

The role of TBBx in fibrotic ILD is further limited by the risk of significant bleeding (1%-4% of cases) and pneumothorax (9%), with half of the latter requiring chest tube drainage.44-50 Based on the limited diagnostic utility and potential risks, a previous consensus statement recommended against TBBx in patients with suspected IPF,3 suggesting that it primarily be used to diagnose infection or malignancy in appropriate settings.

Key messages: Transbronchial Lung Biopsy

- Transbronchial biopsy can be useful in the evaluation of some nonfibrotic ILD subtypes (eg, sarcoidosis) and to identify infection or malignancy.
- Transbronchial biopsy is associated with an increased risk of complications, including bleeding (1%-4%) and pneumothorax (9%) in patients with fibrotic ILD.
- Due to a limited diagnostic yield and increased risk of complications, transbronchial biopsy should not be used routinely in the evaluation of patients with fibrotic ILD.

Surgical lung biopsy

When should surgical lung biopsy be performed in the evaluation of fibrotic ILD?

Surgical lung biopsy (SLBx) is performed using single-lung ventilation under general anesthesia. SLBx was historically performed via open thoracotomy (ie, "open lung biopsy"), but is now typically performed using video-assisted thoracoscopic surgery (VATS), a technique associated with lower perioperative morbidity and mortality.51,52

SLBx for histopathologic sampling in the evaluation of fibrotic ILD should be considered when a definitive ILD diagnosis cannot be established using noninvasive data. A recent systematic review of 2148 patients from 23 studies showed SLBx had a median diagnostic yield of 95% (range 42–100) and changed management in 42%-90% of cases in the 8 studies that reported longitudinal data.52 Two studies reported yields below 70%, although these outliers were limited by the lack of SLBx review in a MDD.53,54

The yield of SLBx can be optimized through several means.3 Biopsies should have a depth ≥2 cm and maximum diameter ≥4 cm when gently inflated post-resection.3,55

Clinicians should clarify the optimal sampling locations with the surgeon as yield is highest for biopsies taken from more involved areas on HRCT, recognizing that areas with severe fibrosis and honeycombing frequently show nonspecific findings and should therefore be avoided.56 Approximately 20% of patients have discordant pathology patterns (eg, UIP in one lobe and NSIP in another lobe),55,57 indicating the importance of sampling at least 2 lobes.3,55,56 In two previous studies, the clinical behavior and prognosis of patients with discordant NSIP and UIP findings were similar to patients with UIP on both biopsies, emphasizing the need to identify a UIP pattern when present.55,57

SLBx for histopathologic sampling in the evaluation of fibrotic ILD should be pursued following a detailed discussion with the patient regarding potential benefits and risks. A recent systematic review and meta-analysis of 1319 patients from 16 studies showed pooled 30- and 90-day post-operative mortality of 2.2% (95%CI 1.0-4.0) and 3.4% (95%CI 1.8-5.5), respectively;52 however, there was significant heterogeneity among studies. A recent retrospective analysis of over 32 000 patients undergoing SLBx showed an in-hospital mortality of 6.4% with significantly higher risk for nonselective compared to elective procedures (16.0% vs. 1.7%). SLBx has been associated with a very low risk of mortality in some cohorts, seemingly related to appropriate patient selection.59 Nonfatal post-operative complications are reported in 8.4%-56% of patients undergoing SLBx, including acute exacerbation of ILD; pneumonia; pleural effusion; chronic chest pain; prolonged air leak; post-operative need for mechanical ventilation; and readmission to hospital within 1 month of discharge.40,58,60-62 Common risk factors and relative contraindications to SLBx are listed in Table 1.52,59,62

Key messages: Surgical Lung Biopsy

- Surgical lung biopsy is a relatively safe procedure with a high diagnostic yield and should be considered when a definitive ILD cannot be diagnosed after a multidisciplinary discussion that incorporates all available data.
- Patients with advanced/rapidly progressive lung disease, significant co-morbidities or who are mechanically ventilated or immune suppressed at the time of surgical lung biopsy are at highest risk of perioperative complications.
- The yield of surgical lung biopsy should be optimized through several means, including:
  1) a biopsy depth of ≥ 2 cm and maximum diameter ≥ 4 cm (when gently inflated post-resection);
  2) sampling of the more severely affected areas on HRCT;
  3) avoidance of areas with severe fibrosis and honeycombing;
  4) sampling of ≥ 2 lobes;
  5) review by an lung pathologist with ILD experience; and
  6) review in a multidisciplinary discussion.

Transbronchial lung cryobiopsy

When should transbronchial lung cryobiopsy be performed in the evaluation of fibrotic ILD?

Transbronchial lung cryobiopsy (TBLC) has been proposed as a histopathologic sampling technique in the evaluation of ILD that potentially provides a balance between the higher yield of SLBx and the lower complication rate of TBBx. TBLC is based on the Joule-Thompson effect, whereby a
compressed gas released at high flow expands quickly, leading to a very low temperature at the tip of a probe passed through the working channel of a bronchoscope. This rapid freezing preserves lung architecture to provide samples with a cross-sectional area of 4–64 mm² compared to ∼1–3 mm² in TBBx. Both cryoprobe and bronchoscope must be removed simultaneously as each biopsy exceeds the size of the bronchoscope’s working channel and patients typically undergo endotracheal intubation for the procedure.

The diagnostic yield and complication rates of TBLC for ILD have been reported in 10 publications from 8 distinct cohorts, although there was significant heterogeneity among studies (eg, patient population, method of sedation, flexible vs. rigid bronchoscope, method of determining the diagnosis). The reported diagnostic yield ranges from 51%-98% when TBLC is guided by HRCT and real-time fluoroscopy, varying by whether the diagnostic yield is reported based on pathological findings assessed in isolation or whether biopsies were reviewed in a MDD. In the only prospective randomized controlled trial, 77 patients with suspected acute or chronic ILD were randomized to either conventional TBBx or TBLC. The TBLC group had a higher percentage of definitive histologic pattern established compared to TBBx (74.4% vs. 34.1%) and a higher percentage of confident MDD diagnoses (51.4% vs. 29.1%). Confident MDD diagnoses were achieved less often than confident histologic diagnoses, which is likely due to discrepancies among clinical, radiological and histological findings, indicating the importance for review of TBLC findings in the context of a MDD. There were no MDD diagnoses of IPF in this study, and it is unknown whether IPF was under-represented in this cohort, or whether the absence of IPF reflects a lack of sensitivity of TBLC for a UIP pattern. Conversely, IPF was the most frequent MDD diagnosis after TBLC in a recent retrospective study of TBLC and SLBx in fibrotic ILD.

There is significant heterogeneity in the reported complications following TBLC with pneumothorax rates ranging from 0%-33% and moderate to severe bleeding ranging from 0%-78%. It is unclear whether this heterogeneity reflects differences in study populations, reporting standards, procedural techniques and/or operator expertise. A prospective study is needed to directly compare the yield and safety of TBLCL to SLBx in the evaluation of fibrotic ILD prior to widespread adoption of TBLC into clinical practice.

### Table 1. Relative contraindications to surgical lung biopsy.

- Age >75 years
- Pre-operative resting hypoxemia
- Mechanical ventilation
- FVC <55% predicted
- DLCO <35% predicted
- Pulmonary hypertension
- Immunocompromised state
- Clinically significant co-morbidity (e.g., ischemic heart disease)
- Rapidly progressive disease

*Note. Abbreviations: FVC, forced vital capacity; DLCO, diffusion capacity for carbon monoxide.*

**Key Messages: Transbronchial Lung Cryobiopsy**

- Transbronchial lung cryobiopsy is a potential approach for histologic sampling of the lung in patients with fibrotic ILD.
- No studies have directly compared the diagnostic accuracy and safety of transbronchial lung cryobiopsy to surgical lung biopsy.
- Additional data are required prior to adoption of transbronchial lung cryobiopsy into clinical practice.

**Multidisciplinary discussion**

Is multidisciplinary discussion required prior to initiation of antifibrotic or immunosuppressive pharmacotherapies in patients with fibrotic ILD?

MDD is the interactive dynamic dialogue among clinicians, radiologists and pathologists with expertise in ILD, with the goal of achieving a consensus diagnosis in patients with suspected ILD. MDD was first recommended in the 2002 consensus statement describing the classification of the IIPs and has since been considered the gold standard for ILD diagnosis. MDD improves diagnostic accuracy (measured as inter-observer agreement) in both community and academic settings compared to diagnoses made independently. Inter-observer agreement is higher among academic compared to community physicians, with the latter group more likely to assign a diagnosis of IPF, illustrating the importance of diagnosing ILD using a MDD among ILD experts. Inter-MDD agreement is acceptable overall for establishing a specific IIP diagnosis, and good for IPF. MDD leads to greater confidence in a diagnosis of IPF, compared to a diagnosis established by individual clinicians or radiologists and may help avoid an unnecessary surgical lung biopsy in some patients.

An accurate ILD diagnosis is particularly important in the context of IPF-specific medications that are not indicated in other fibrotic ILDs and considering the harm caused by inappropriate use of immunosuppressive medications in IPF. The strong rationale and improved diagnostic accuracy have prompted multiple groups to recommend that ILD diagnoses be established via MDD whenever possible, including review of at least clinical, laboratory and radiological data. Patients that demonstrate unanticipated clinical changes or that subsequently undergo SLBx should be re-presented, illustrating the iterative nature of a MDD. A substantial proportion of ILD patients cannot be confidently classified with a specific ILD diagnosis despite detailed review of all available data; however, a MDD can often help guide management decisions even in the absence of a confident IIP diagnosis. Patients with “unclassifiable ILD” should periodically be re-evaluated by an ILD expert and MDD group to evaluate new findings that may suggest a specific underlying diagnosis. This general approach to ILD diagnosis is summarized in Figure 1, highlighting the iterative nature of the MDD with a proposed schematic framework to the evaluation of fibrotic ILD.

Access to a formal MDD is frequently restricted by time constraints, geographic barriers, lack of remuneration for participating physicians and the limited number of experienced ILD centres. Some Canadian ILD centres offer review of ILD cases through tele-health or other similar means, although it is unknown how this process compares to a standard MDD. Despite these potential barriers, it is appropriate to review ILD patients in a MDD setting whenever possible before initiating...
disease-specific pharmacotherapies, considering the costs and potential adverse effects of these medications.

Key Messages: Multidisciplinary Discussion

- Multidisciplinary discussion is the interactive dynamic dialogue among expert respirologists, radiologists and pathologists that is currently considered the gold standard for ILD diagnosis.
- A multidisciplinary discussion is an iterative process and patients should be re-reviewed if new patient information becomes available after the initial discussion (eg, new histopathological results or a change in radiological pattern).
- ILD patients should be reviewed in a multidisciplinary discussion whenever possible before initiating disease-specific pharmacotherapies.

Knowledge transfer and tools for practice

- The present document is available for download at www.respiratoryguidelines.ca and www.tandfonline.com.
- A slide deck for teaching and self-learning as well as a handout for health care professionals and students is available at www.respiratoryguidelines.ca.
- The CTS Clinical Assembly on Interstitial Lung Disease welcomes the opportunity to partner with other organizations and stakeholders in the development of educational tools and resources that support the implementation of the key messages described herein, with various targeted groups.
- Successful implementation of the clinical guidance in this position paper is integral to its aims, and this may be monitored through surveys of document end-users, as well as administrative database analyses of diagnostic testing in the evaluation of patients with suspected fibrotic ILD.

Conclusions and future directions

Fibrotic ILD represents a large heterogeneous group of disorders that are challenging to diagnose. The evaluation of patients with fibrotic ILD should include a thorough history, physical examination, basic serological testing, pulmonary function tests and HRCT of the chest. Specific serologic testing, BAL, TBBx and SLBx may be informative in select patients depending on the clinical presentation and diagnoses under consideration. Where feasible, fibrotic ILD cases should be reviewed in a MDD that includes clinicians, radiologists and pathologists with expertise in ILD. This comprehensive patient evaluation aims to provide adequate clinical data to enable accurate diagnoses, consistent with disease-specific diagnostic criteria and a standardized ontological framework. There is a relative paucity of data to guide these recommendations in the broad category of fibrotic ILD and the key messages in this document represent the consensus opinion of ILD experts from a diversity of disciplines across Canada. In certain clinical contexts, there may be barriers to implementing these recommendations, including limited clinical experience or access to specific clinical resources. Future work should identify these barriers while testing the efficacy of specific interventions to
improve the implementation of evidence-based recommendations. Further research is needed for each modality previously outlined to inform their performance characteristics and clinical utility in the evaluation of fibrotic ILD. We also note that we did not consider the views and preferences of the target population (patients with fibrotic ILD) in this consensus statement development process, and these valuable insights should be sought in future position statements or clinical practice guidelines.

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Declaration of interest

The CTS Clinical Assembly on Interstitial Lung Disease is accountable to the CTS Respiratory Guidelines Committee and the CTS Board of Directors. The CTS Clinical Assembly on Interstitial Lung Disease is functionally and editorially independent from any funding sources of the CTS and does not receive any direct funding from external sources. The CTS receives unrestricted grants that are combined into a central operating account to facilitate the knowledge translation activities of the CTS Clinical Assemblies. No funders played a role in the collection, review, analysis or interpretation of the scientific literature or in any decisions regarding the key messages presented in this document.

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