Introduction

Interstitial lung diseases (ILDs) are the most common pulmonary manifestation of connective tissue diseases (CTDs) and are often the most serious complication. Lung involvement in CTDs is associated with decreased survival rates and higher overall mortality. ILD may occur in almost all clinically diagnosed CTDs, but it is more prevalent in systemic sclerosis (SSc) and idiopathic inflammatory myopathies (IIM). The specific type of ILD can be characterized based on diverse patterns of lung damage seen on high resolution computed tomography (HRCT) and by histopathological analysis of lung biopsy specimens. These diverse forms of ILD were discussed in previous chapter; they mainly comprise of the following types: usual interstitial pneumonia (UIP), nonspecific interstitial pneumonia (NSIP), diffuse alveolar damage pattern (DAD), lymphocytic interstitial pneumonia (LIP) and organizing pneumonia (OP). Management and treatment options for ILD-CTD have been extensively studied, but they still pose a significant challenge due to the paucity of randomized controlled trials (RCTs). It should also be noted that there can be other pulmonary manifestations of CTDs (other than just ILD): these include pleural effusions and bronchiectasis seen with systemic lupus erythematosus (SLE) and rheumatoid arthritis (RA). These effusions are usually bilateral, secondary to pleural inflammation and often resolve with treatment of underlying CTD.

This chapter will describe the various treatment options for ILD-CTDs as described by medical literature while also reviewing ongoing clinical trials. We will go through treatment options for ILDs based on the specific type of CTD.

Disease Specific Treatment

Systemic Sclerosis (SSc)

Systemic Sclerosis involves a complex pathogenesis of immune system activation, vascular constriction and increased activity of fibroblasts. The resultant fibrosis can affect every organ of the body and lungs are no exception. ILD is the most prevalent form of pulmonary involvement, and NSIP is the most implicated pattern.

Precise criteria for initiation of immunosuppressive regimen for systemic sclerosis-associated interstitial lung disease (SSc-ILD) has not been established yet. The selection of treatment requires knowledge of its toxicity while providing it to those whose disease...
is most likely to progress. In progressive SSc-ILD, the most rapid decline in functional vital capacity (FVC) often occurs within first three years of disease onset and thus timing of therapy becomes crucial in such cases13.

Approach to Therapy

HRCT and pulmonary function test (PFT) results play an important role in determining the timing of initial therapy. Serial PFTs that demonstrate 10% decline in FVC or 15% decline in diffusing capacity for carbon monoxide (DLCO) are indications of active disease for which treatment should be considered14. Similarly, HRCT demonstrating fibrosis of 20% or more of lung parenchyma has been associated with increased mortality and thus should guide towards early therapy initiation15,16. Additionally, SADL Model (smoking history, Age and % predicted DLCO) has been shown to estimate 3-year mortality risk in SSc-ILD using readily available clinical parameters and thus can be utilized for therapy initiation conversation17.

Initial Therapy

Mycophenolate mofetil (MMF) and Cyclophosphamide (Cyc) have both been used and studied in multiple observational studies and RCTs as initial therapies for SSc-ILD. MMF has been shown to decrease the decline of DLCO, stabilize decrease in FVC and improve respiratory symptoms such as dyspnea and cough10,16.

Scleroderma Lung Study II (SLS II) trial showed that the difference of improvement in FVC between MMF and oral Cyc was not significant. Cyc was however associated with higher rate of adverse events such as leukopenia and thrombocytopenia20, and thus MMF is often deemed a superior initial agent due to safer profile. On the contrary, MMF can cause gastrointestinal side effects such as cramping, diarrhea and nausea, this may be abated by using enteric coated formulations and dividing dosing regimen.

Cyclophosphamide is also a first line agent used for progressive SSc-ILD and has shown to improve fibrosis and FVC31,22. Both oral and IV Cyc can be used but latter is preferred due to a better side effect profile29. Cyc usage is mainly limited due to its adverse effects of hemorrhagic cystitis, leukopenia, and thrombocytopenia. If utilized, its duration is limited to 6-12 months with switching to a less toxic agent for maintenance thereafter.

Low dose glucocorticoid (GC) (<10mg/day) has been used with Cyc (in addition to other immunosuppressants) in studies demonstrating Cyc's efficacy, however evidence for its usage is scarce and long term therapy is precluded due to adverse effects related with GC. High dose glucocorticoid are avoided with Cyc due to increased risk of propagating scleroderma renal crisis31.

Maintenance Therapy

Both MMF and Azathioprine (AZA) can be administered for maintenance therapy after an initial round of MMF/Cyc25,26. MMF is more widely used though due to its proven benefit and efficacy20. These agents are utilized until patient experiences a period of clinical stability in respiratory symptoms or adverse effects warrant use of a different therapeutic option. However, the optimal duration of maintenance therapy is yet to be known.

Newer Options

Elevated levels of Interleukin-6 (IL-6) are associated with progression of SSc-ILD27, and earlier in 2021 FDA approved use of tocilizumab in such patients to halt the rate of decline of lung function28. Tocilizumab has been shown to decrease rate of FVC change as compared to placebo in clinical trials29,30, but there have been no comparisons with MMF directly and thus its usage depends on limitation of MMF or Cyc therapies due to side effects.

Nintedanib is an FDA approved tyrosine kinase inhibitor for idiopathic pulmonary fibrosis(IPF)31, it has also been studied in patients with SSc-ILD where it showed considerable improvement in rate of decline of FVC32, however more RCTs need to be undertaken to better assess its efficacy. Nintedanib was approved by the FDA and EMA recently for patients with SSc-ILD and chronic fibrosing ILD (based on results of the SCENSIS and INBUILD trial). Pirfenidone is commonly used to slow progression of IPF, and it has shown improvement in FVC in SSc-ILD patients as well33,34. However, larger studies with longer follow-up periods are required for conclusive evidence.

Rituximab has also been studied in a few observational studies and small trials. It shows promising results with improvement in FVC and DLCO compared to standard therapy35-37, however its use is limited owing to lack of larger RCTs.

Idiopathic Inflammatory Myopathies (IIM)

IIM comprise of Dermatomyositis (DM), Polymyositis (PM) and Antisynthetase Syndrome (ASSD). These CTDs involve varying degrees of skeletal muscle inflammation with frequent extra-muscular organ involvement. ILD is the most common extra-muscular manifestation of IIM with NSIP being the most prevalent form38, while OP is also frequently seen with IIM.

Approach to Therapy

As discussed earlier with SSC-ILD, therapeutic approach for IIM-ILD depends on degrees of respiratory symptoms (dyspnea),PFT results, degree and type of fibrosis on HRCT and histopathology, and progression of ILD39. It is seen that patients who are positive for melanoma differentiation-associated (MDA) – 5 antibodies have worse survival rates
compared to those who don’t, thus necessitating earlier treatment. On the contrary, anti-aminoacyl tRNA synthetase antibodies, a group of myositis-specific autoantibody related to ASSD are associated with a better glucocorticoid response.

**Initial Therapy**

Systemic Glucocorticoids form the mainstay of initial therapy for IIM-ILD. Even though there has been no controlled trial demonstrating its efficacy, anecdotal usage has shown that almost half of the patients with DM or PM associated ILD clinically improve with GC. There are also certain clinical/laboratory features that propose high response to GC therapy; younger age, elevated creatine phosphokinase (CPK) levels, PM etiology instead of DM, NSIP rather than UIP on histopathology and ground glass opacities or consolidation on HRCT compared to honeycombing.

A steroid sparing agent is usually used with initial GC therapy regardless of response to GC.

**Second Agent Options**

Exact guidelines to start a second steroid sparing agent have not been yet established but certain clinical features are strong indicators to start a second agent therapy: Impending respiratory failure, antisynthetase syndrome with lung involvement or anti-MDA 5 seropositivity as these are associated with rapidly progressive ILD.

MMF, AZA, Methotrexate (MTX) and Calcineurin inhibitors have all been used as steroid sparing agents, their choice depends on toxicity of the treatment, agent being used to treat myositis and severity of lung involvement. Both AZA and MMF show comparable efficacies with improvement in HRCT picture, FVC and DLCO, however larger studies are still required to further evaluate the roles of these medications. Tacrolimus and Cyclosporine administration have shown improvement in survival and lung parameters in a few studies, however lack of larger trials and risk of nephrotoxicity limit their usage to refractory cases. MTX is commonly used as a steroid sparing agent in IIM, however it may cause drug-induced pneumonitis, which may be difficult to differentiate from underlying ILD. Cyclophosphamide may be used for more severe and refractory cases of ILD.

Rituximab has been shown to improve PFT parameters (DLCO and FVC) in patients with refractory myositis-related ILD (especially ASSD). This association is limited to a few case reports/series.

**MDA-5 Antibody Seropositivity**

As previously mentioned, MDA-5 Antibody seropositivity portends a rapidly progressive ILD refractory to Cyc, GC and calcineurin inhibitors. Rituximab and Janus Kinase (JAK) inhibitor tofacitinib have been shown to improve ILD in these patients. In an open-label study, tofacitinib usage demonstrated 100% survival at 6-month interval compared to other immunosuppressants (GC, Cyc, MMF, AZA) in this patient subset. It was also linked with improvement in DLCO and HRCT findings. Similarly, plasma exchange and IVIG administration has shown improvement in this subset of patient population, but larger trials are required to better evaluate efficacy of these options.

**Rheumatoid Arthritis**

Rheumatoid arthritis has many extra-articular manifestations and lungs are commonly affected; with ILD being the most common form of pulmonary disease. Rheumatoid arthritis related Interstitial lung disease (RA-ILD) is a heterogeneous group of different subtypes of ILDs with differing clinical approaches and treatment options. UIP and NSIP are often the most implicated histopathologic and radiographic pattern in RA-ILD.

**Approach to Therapy**

RA-ILD’s treatment modalities continue to be researched on and no optimum therapy guideline is available due to a severe lack of RCTs in this subset of patients. The decision to start immunosuppressive therapy is undertaken after assessment of prognosis, adverse effect of treatment and rate of progression of disease. A pragmatic approach usually involves characterizing disease behavior into categories such as self-limited, reversible, progressive, or irreversible based on serial measurement in predictors of survivals such as DLCO and HRCT picture.

**Glucocorticoid (GC)**

GCs are commonly utilized for initial RA-ILD therapy. Certain features pertain a more favorable response to GCs including younger age, non-UIP histopathologic disease and worsening of HRCT and/or pulmonary function.

If patients fail to show improvement in lung function on GC therapy, an approach involving other immunosuppressants such as MMF, CYC and AZA can be considered after excluding drug toxicity and infection as cause of failure to respond. These agents can also be utilized as GC-sparing regimen after initial therapy with GC or in conjugation with them as well in order to minimize adverse effects related with GC.

**Methotrexate (MTX) Therapy**

MTX has historically been linked with worsening of pulmonary fibrosis, but newer data suggests that MTX may in fact delay the onset of RA-ILD. Thus, clinically MTX therapy is used if indicated for underlying joint disease.

**Progressive Fibrosing ILD**

Patients who have >10% lung involvement in fibrosis
or have feature of extensive reticulation with traction bronchiectasis are defined as having progressive fibrosing ILD. This form of ILD is a broader term which can be seen with other CTD-ILDs and is characterized by progressive decline in lung function (decline in FVC/DLCO >10%) despite appropriate therapy. Nintedanib has been studied in RA-ILD patients with progressive fibrosis and shows improvement and stability in lung function71. However, lack of individualized trials on just RA-ILD patients necessitate continuing research.

Pirfenidone is currently being studied for RA-ILD management in a phase 2 clinical trial72. These results and studies on other newer options will direct the course of future treatment options.

**Newer Options**

Abatacept, Rituximab and Tocilizumab have all shown modest control of RA-ILD in retrospective studies73-76, however there is a need for larger clinical trials to formally assess the efficacy of these treatment options.

**Sjogren’s Syndrome (SS)**

Sjogren’s Syndrome involves lymphocytic infiltration of salivary and lacrimal glands resulting in their destruction. SS related ILD is often the most common manifestation of associated lung disease with NSIP being the most found pattern. UIP is occasionally observed and even though LIP is rarely observed, it is classically associated with SS77.

**Approach to Therapy**

Asymptomatic patients can be monitored over a course of months – years with serial PFTs and HRCT to document any worsening in clinical picture as primary SS-ILD often progresses more slowly. For symptomatic patients with worsening symptoms and/or deteriorating HRCT or PFT parameters treatment is usually indicated78. Though it needs to be noted that treatment options are usually empiric and there are no formal guidelines available specific for SS related ILD treatment as data is limited to a few case reports/series.

**Initial Agent**

GCs are frequently used as initial therapy, response to GCs and GC-sparing agents is usually good with CYC reserved for more severe cases. MMF or AZA can be used to minimize steroid-related side effects and to improve response to GC67,79,80. Patients with radiologic/histopathologic evidence of NSIP, LIP or OP seem to show a better response to immunosuppressive treatment than those who show evidence of a UIP pattern.

**Refractory Disease**

Few case reports and case series have shown improvement and stability in lung function with use of Rituximab81-85 thus this option can be utilized for SS-ILD patients refractory to GC and conventional immunosuppressant therapy, though it should be noted that larger trials are still required.

Tocilizumab usage improved SS-ILD in one case of steroid resistant OP86.

**Systemic Lupus Erythematosus (SLE)**

SLE is an autoimmune disorder characterized by autoantibodies against nuclear antigens and immune complex deposition in various organs. NSIP is the most common ILD pattern found in SLE patients with OP and LIP pattern being occasionally described87. It should be noted that incidence of ILD in SLE is overall relatively low.

Therapeutic options for SLE-ILD are mostly dependent on anecdotes as no clinical trial has been done specifically for this subset. Glucocorticoids are usually the initial agents used88, with MMF and AZA acting as steroid sparing agents89. CYC and Rituximab may be used for cases resistant to conventional treatment options but evidence is limited due to lack of controlled trials90.

**Mixed Connective Tissue Disease (MCTD)**

MCTD is defined as an overlapping condition with clinical features of SSC, SLE, RA, and IIM.

Lung involvement in MCTD is usually like SSc related ILD. There have been no controlled studies assessing treatment options in specifically patient with MCTD.

Similar to treatment of ILD related to CTDs, immunosuppressive therapy can be utilized with combination of GCs with AZA and MMF as steroid sparing options91-93, with CYC reserved for refractory cases.

**Conclusion**

ILD is the most serious manifestation of lung disease associated with CTDs and is often the cause of death in these patients. It imposes a significant clinical challenge to rheumatologists and pulmonologists due to many factors. Some of these factors include lack of large clinical trials demonstrating efficacy of treatment options, unpredictable nature of the disease, lack of clear underlying histopathologic diagnosis and grim prognosis of overall condition.

Therapeutic approach for treatment of CTD-ILD is obtained from mostly retrospective or inconsistent data, apart from approach for SSC related ILD. Therefore, a lot of times clinicians paint every CTD-ILD with the same brush and use similar treatment options. This undermines the diversity in types, pathogenesis, and therapy response among different types of ILD.

Clinically, treatment approach is based on initial
immunosuppression with glucocorticoids and immunosuppressive regimens, mainly CYC, AZA and MMF, forming the steroid sparing group. Even though strong evidence is still not substantiated, Rituximab is considered a strong contender for both initial therapy as well as for refractory disease\footnote{94}. More recent clinical trials have introduced an array of newer options for CTD-ILD treatment including Nintedanib and Tocilizumab amongst others\footnote{94,95}. Having said that, larger controlled trials are still required to formulate treatment guidelines for CTD-ILDs.

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