Transbronchial lung cryobiopsy in the diagnosis of interstitial lung disease

Joseph KEENAN1, Elliot BACKER1, Heidi GIBSON2, Roy CHO1, H. Erhan DINÇER1

1 Division of Pulmonary, Allergy, Critical Care and Sleep Medicine, Minnesota University, Minnesota, USA
1 Minesota Üniversitesi, Solunum, Allerji, Kritik Bakım ve Uyku Bölümü, Amerika Birleşik Devletleri
2 Division of Cardiopulmonary Services, Minnesota University, Minnesota, USA
2 Minesota Üniversitesi, Kardiyopulmoner Bölümü, Amerika Birleşik Devletleri

ABSTRACT

Transbronchial cryoprobe lung biopsy (TBCLB) have recently been introduced as a safe diagnostic tool in the diagnosis of interstitial lung diseases. While we do not enough evidence its role and place as a diagnostic procedure, the technique has been adopted by many centers. In spite of expanding body of literature, there are variations in patient selection and procedural aspect of the procedure. It has been established as a safe procedure if safety measures are practiced. Diagnosis of interstitial lung diseases continuous to be challenging. Surgical lung biopsy considered as gold standard but its morbidity and mortality limit its utilization in every case. Multidisciplinary medical decision is a validated team work effort when approaching patients with interstitial lung disease.

Key words: Cryobiopsy; transbronchial; interstitial lung disease

ÖZ

Interstisyal akciğer hastalığı tansında transbronşiyal akciğer kriyobiyopsi

Transbronşiyal kriyoprob akciğer biyopsisi (TBCLB), son zamanlarda interstisyal akciğer hastalıklarının tansında güvenli bir tani yöntemi olarak gösterilmektedir. Tanıdaki rolü ve yerindeki yerine kant bulunmasına da, teknik birçok merkez tarafından kabul edilmiştir. Literatür verilerinin araştırıma rağmen, hasta seçiminde ve prosedürün uygulanmasında farklılıklar vardır. Cerrahi önlemlerin uygulandığı durumlarda, güvenli bir prosedür olarak belirlenmiştir. Interstisyal akciğer hastalıklarının tansundaki zorluklar devam etmektedir. Cerrahi akciğer biyopsisi altın standart olarak kabul edilmektedir, ancak morbidite ve mortalitesi nedeniyle uygulanması kısıtlıdır. Interstisyal akciğer hastalıkların tansında uygulanması, multidisipliner tibbi karar onaylanması bir ekip çalışmasıdır.

Anahtar kelimeler: Kriyobiyopsi; transbronşiyal; interstisyal akciğer hastalığı
Transbronchial cryoprobe lung biopsy (TBCLB) is a relatively new technique for sampling lung tissue. This technique relies on the fact that rapidly frozen lung tissue sticks to the tip of the probe allowing larger biopsy specimens with less crush artifact (1,2). The diagnostic yield of transbronchial biopsies with standard forceps is often limited due to smaller size and crush artifact. Therefore, TBCLB offers a potential diagnostic benefit over traditional transbronchial biopsies. In recent years, TBCLB has gained popularity and been used various lung conditions, such as interstitial lung diseases (ILD) or diffuse parenchymal lung disease (DPLD), lung transplantation and lung cancer, to improve diagnostic yield (3-6). Overall, traditional forceps transbronchial biopsies play a minor role in the diagnosis of ILD and it has mainly been used to rule out granulomatous diseases (e.g. sarcoidosis) or infection. Surgical lung biopsy is considered as gold standard for the diagnosis of ILD largely because of large parenchymal tissue allowing evaluation of larger area of preserved architecture and subpleural tissue. Unfortunately, many patients cannot undergo surgical lung biopsies due to underlying advanced lung disease. Moreover, possibility of ILD exacerbation makes clinicians not to choose surgical biopsy as first option. Since the first feasibility study on safety and yield of TBCLB performed by Babiak in 2009, many other studies, mostly retrospective, have been published last several years (5). There are still no widely accepted guidelines on biopsy technique, patient selection, number of biopsies, tissue processing, airway management and anesthesia. This review intends to provide information on diagnostic yield, biopsy techniques and potential complications.

**Definition of DPLD and Diagnostic Modalities**

DPLD or ILD is a group of diseases mainly affecting the all three components of the lung; endothelium, interstitium and epithelium. DPLD encompasses over 140 different diseases that eventually lead to increasing cellularity and fibrosis. Prognosis and treatment options vary based on underlying cause, if known. Patients may have a range of outcome from complete remission with or without treatment to progressive and fulminant course. The current recommendation indicates that a multidisciplinary expert team of pulmonologists, radiologists and pathologists should be involved in a consensus clinical diagnosis (7,8). Initial step in diagnosis is taking a thorough history that includes domestic and occupational environmental exposures, connective tissue disease (CTD) or drug toxicity for DPLD and a family history for familial form of pulmonary fibrosis. If a potential cause is identified, the patient should be evaluated to rule in or rule out other known causes such as CTD, hypersensitivity pneumonitis (HP), pneumoconiosis and iatrogenic causes (e.g. drugs and irradiation). If a specific diagnosis or potential cause is not identified, further work up should be done by serology and blood work for CTD and HP and high-resolution CT scan of the chest to identify the pattern of involvement. HRCT may reveal a specific pattern leading to a specific diagnosis such as usual interstitial pneumonia (UIP). If the changes suggest DPLD, a multidisciplinary discussion is strongly suggested to determine the need of an invasive procedure. Surgical lung biopsy is considered as gold standard but surgical biopsy in some of the HRCT patterns such as UIP or some familial cases is not necessary. Moreover, surgical biopsy would not be feasible in patients at high risk for respiratory failure, exacerbation of DPLD or prolonged air leak, especially those with hypoxemia at rest, severe pulmonary hypertension or a corrected diffusion capacity of carbon monoxide for hematocrit less than 25% of predicted (9).

Among invasive procedures, bronchoalveolar lavage is not routinely recommended however can be appropriate when the radiologic differential diagnosis include eosinophilic pneumonia, sarcoidosis or infection. Cell counts (e.g. higher neutrophil proportion in interstitial pulmonary fibrosis, lower proportion of alveolar macrophages in IPF, higher eosinophils in eosinophilic pneumonia and higher CD4/CD8 in IPF and sarcoidosis (10-12).

Transbronchial forceps lung biopsies (TBFLB) has been utilized in attempt to make diagnosis in DPLD however, there are no studies about clinical outcomes in those who underwent TBFLB or not. Current guidelines do not recommend performing TBFLB in patients who are clinically suspected of having IPF with an UIP pattern on HRCT. However, there is no recommendation for or against performing TBFLB in patients with clinically suspected IPF and HRCT findings of probable or intermediate IPF. Nevertheless, unweighted studies indicated that almost three-fourth of cases resulted in adequate sampling and half of those with adequate sampling led to a definitive diagnosis eliminating the need of surgical biopsy (13-15).

Although surgical lung biopsy (SLB) is considered as gold-standard due to the fact that an adequate tissue can be sampled in almost all cases where paucity and
heterogeneity of parenchymal involvement can be captured as well as subpleural tissue that is important in the diagnosis of UIP. SLB may not be an option in some patients with comorbidities and may result in post-operative complications such as prolonged air leak, bleeding, infection and ILD exacerbation. Risk of procedural mortality, although rare, does exist. Based on most studies, diagnostic yield of SLB was 88%-90%. Although there is role in the diagnosis of ILD for SLB, especially when the multidisciplinary discussion is the decision-maker, SLB is not recommended in patients with clinical suspicion of IPF and HRCT pattern of UIP (8,16-19).

Role of TBCLB in the diagnosis of DPLD has been gaining popularity although there is no study compared clinical outcomes for those underwent to biopsy or not. Pooling studies indicated an adequate sampling in majority of cases (94% to 97%) in which a definitive diagnosis was achieved in 77%-83%. TBCLB is considered to be more invasive than TBFLB and reported have complications of death, bleeding, pneumothorax and prolonged air leak, although the incidence of procedure related death is less than SLB, 0.04%-1.3% vs. 0.8%-3.5%, respectively. Based on current guidelines, TBCLB is not recommended in patients with clinically suspected IPF and HRCT pattern of UIP while no specific recommendation was given for those with probable UIP or indeterminate UIP changes on HRCT (8,20-23).

Cryothechnology and its Application in Pulmonary Medicine

Currently, there are rigid, semi-rigid and flexible cryoprobe lung biopsies with tip diameters of 1.9 mm, 2.4 mm and 5.5 mm and length of 90 cm, 50 cm and 60 cm (Spembly Medical Ltd, UK and ERBE, Germany). Cryo equipment has 2 components in addition to the probe; 1. the console that monitors the cryopressure, probe tip temperature and the length of procedure and 2. Cryogen such as nitrous oxide, carbon dioxide or liquid nitrogen to free the tip of the probe to -70°C, -89°C and -196°C, respectively. The technology relies on Joule-Thompson effect in which compressed cryogen gas exits at a high flow, expands rapidly and creates low temperatures to allow tissue adhesion to the tip of the probe (24). Cryotechnology has been used in pulmonary medicine for ablation and diagnostic purposes. Ablation of benign or malignant lesions with cryotherapy relies on the destructive effects that are 2-fold: cellular injury and vascular injury. Ablation effect of cryo technology is beyond the scope of this review therefore we will not review it further.

Cryoprobe Lung Biopsy Techniques

In 2009, Babiak et al. has introduced cryoprobe lung biopsies as a new tool (5). The biopsy technique has many variations among centers worldwide. Simply, the probe is advanced through the working channel of the bronchoscope into the lung periphery and then the probe is activated with a foot pedal for few seconds causing an ice-ball around the tip of the probe where the lung parenchymal tissue adheres to the ice-ball and probe. At this point, the bronchoscope and the cryo probe attached lung tissue are retracted en-bloc from the airway. Then, the probe is either submerged into a saline or a saline stream instilled on to the frozen probe tip with lung tissue to separate the lung tissue specimen from the probe. Technical variations derive from operators training and logistics within their institution. However, the most important factor plays role in variation is the readiness for dealing with potential complication of the procedure. The potential complications encountered in TBCLB are hemorrhage that maybe excessive leading to respiratory compromise and pneumothorax. It should be remembered that some of the ILD patients have limited respiratory reserve making them vulnerable to complications. Besides dealing with complications, operators may select different techniques when acquiring biopsies that involves selection of probe size (1.9 mm vs. 2.4 mm), duration of freezing time, radial EBUS guidance, biopsy location (central or peripheral or pleural based) and number of biopsies in one or multiple lobes (1,3-5,23,25-29).

At present time, there is no single technique that is accepted by the operators. Our group performs cryo probe lung biopsies mostly in the OR while the patient is intubated with an ET'T of 8.5 mm and under general anesthesia. We target the areas of mostly ground-glass opacifications and at least 2 cm away from the pleural surface. The biopsies (3 to 5 biopsies from each lobe) are taken under live fluoroscopy with potential complication of the procedure. The duration of freezing is decided based on the first biopsy size that is usually taken at 4 second mark of freezing time. If the biopsy size is less than 3 mm we increase the freezing time 2 to 4 seconds. We do not use 2 bronchoscope technique or routine balloon occlusion after each biopsy (25,30).
Patient Selection and Safety Concerns

Patient selection is important to alleviate potential procedural complications. There are currently no guidelines about pre-procedural evaluation.

Anticoagulation and antiplatelet agents should be held appropriate time based on their biological half-lives. It is generally an accepted practice to keep international normalized ratio (INR) lower than 1.5 and thrombocyte count above 50,000/µL (31-33). However, it should be noted that these recommendations are made for transbronchial forceps biopsies or EBUS guided needle biopsies. There are no trials about safety of cryoprobe lung biopsies in patients with coagulopathy or thrombocytopenia. In our practice, we prefer to keep INR less than 1.5 and platelet count above 100,000/µL.

Uremia is known to cause platelet dysfunction and increased bleeding risk that is shown to be alleviated with desmopressin (DDAVP) infusion prior to biopsies, risk of bleeding reduced from 45% to 4% (34-36). Although no evidence, we give DDAVP in those with BUN or greater than 35 mg/dL.

Pulmonary hypertension is a common complication of ILD and defined as a resting mean pulmonary artery pressure equal or greater than 25 mmHg. Although right heart catheterization is considered as gold-standard for the diagnosis of pulmonary hypertension, we prefer to screen ILD patients with trans-thoracic echocardiogram prior to cryoprobe biopsies. Although there is no evidence we do not perform cryoprobe lung biopsies if the mean pulmonary arterial pressure is greater than 35 mmHg (those with moderate pulmonary hypertension). It should be remembered that preprocedural echocardiogram has not universally obtained in most of the studies involving cryoprobe lung biopsies.

Similarly, pulmonary function testing has been used to exclude some of the patients from performing cryoprobe lung biopsies, those with FEV1 < 0.8 L or < 50% of predicted, FVC < 50% predicted and DLCO < 35% or 50% of predicted, albeit a recent meta-analysis of 994 cryobiopsy cases did not find a adverse relationship between poor pulmonary functions and complication rates (28,37,38).

Significant hypoxemia (PaO2 less than 55 mmHg on room air) is also considered as contraindication in some of the series (28,37,38).

Complications

Main complications from cryoprobe lung biopsies consists of bleeding, pneumothorax and ILD exacerbation.

In spite of precautions exercised against coagulation, bleeding parameters, medications and pulmonary hypertension in terms of patient selection and pre-procedural patient preparation, bleeding complication has been reported common after cryoprobe biopsies. Unfortunately, there is no accepted scale of bleeding and difficulty to measure bleeding amount, it is hard to compare the incidence of severe bleeding. Most publications utilize a 4-step bleeding as measurement; no bleeding, mild bleeding requiring suctioning only, moderate bleeding requiring endoscopic interventions (e.g. instillation of cold saline, balloon occlusion/tamponade) and severe bleeding leading to respiratory or hemodynamic instability. A meta-analysis of 12 studies and 383 patients, 16.9% reported to have moderate bleeding with a pooled probability of 0.12 (95% CI 0.02-0.25) (40). Although there are no reports of bleeding related mortality in the medical literature, operators should be ready to act on life-threatening bleeding that is a possibility. In our experience, we encounter more bleeding when biopsies are taken from a more central lesions than peripheral lesions that can be explained by the fact that pulmonary artery diameter is larger in adjacent to the central bronchi. Routine usage of bronchial blocker after each biopsy is not standard of care but performed by some centers. However, we strongly recommend having bronchial blockers readily available if it is not used after each biopsy. Although this is anecdotal, we encounter more bleeding in transplant patients likely secondary to nerve denervation leading to poor vasospasm.

The rate of pneumothorax is variably reported in studies. The incidence was reported as 10% and 9.5% in 2 recent meta-analyses (40,41). It is believed that UIP histology, biopsies of fibrotic areas and biopsies close pleural surface increase the risk of pneumothorax risk (37,40). Unfortunately, most studies did not provide information about the severity and duration of bronchopleural fistula. As a precautionary measure, we always keep different sizes of chest tubes in the procedure room, examine the chest with ultrasound just after and a chest X-ray one hour after the procedure.

Acute exacerbation of ILD (AE-IPF) is best characterized in IPF but can occur in other fibrotic ILDs.
Diagnostic criteria for AE-IPF has been updated in 2016 (42). AE-IPF has been described after video-assisted thoracoscopic wedge resection, lobectomy, surgical open lung biopsy and after bronchoalveolar lavage. There are several reports of acute exacerbation of IPF after cryoprobe lung biopsy with incidence up to 2.3%. Although this is not a common occurrence, operators should be aware of this complication and discuss with patients prior to procedure (43-45).

It is also unclear if the centers track their 30-day and 90-day mortality rates following TBCLB. Although meta-analyses report procedural mortality rates of 0.1-0.3%, a retrospective study of 197 patients whom underwent cryoprobe biopsies noted 2% 30-day and 2.5% 90-day mortality. All patients died within 30 days of biopsy had poor baseline lung function of less than 35% predicted DLCO value (46).

**Diagnostic Yield of Cryoprobe Lung Biopsy**

While the role of TBCLB in the diagnosis of interstitial lung diseases is evolving, it remains to be an option as a diagnostic tool. Although there are no clear-cut indication criteria about when to perform TBCLB in the diagnosis of ILD, an expert team including intervention pulmonologist should be part of the decision-making process. Currently, cryoprobe lung biopsy is considered when SLB is considered. Some centers favor TBCLB as an initial invasive diagnostic procedure and consider SLB if no definitive diagnosis is established. This is based on the fact that SLB may result in higher morbidity and mortality than cryoprobe biopsies. It is reported that AE-IPF is more common and mortality may occur after SLB (44,47,48).

Diagnostic yield of TBCLB reported > 50.6% to 100% in various studies. It should be noted that multidisciplinary discussion (MDD) should weigh in final diagnosis and histologic evaluation should be performed by an experienced lung pathologist (49). In our opinion, many factors influence the final diagnosis; MDD, experience of lung pathologist, size and number of biopsy specimens, presence and amount of alveolar tissue and location of biopsies (subpleural, areas of ground-glass opacifications). TBCLB has been utilized in patients for DPLD but very few studies have published on diagnostic yield of cryoprobe lung biopsy exclusively in patients with ILD. As example, our group published a series of 40 ILD patients in whom TBCLB established a confident histopathologic and MDD consensus diagnosis in 85% of the cases (30).

A recent study reported a poor concordance between sequential TBCLB and SLB in the diagnosis of diffuse interstitial lung disease. 21 patients initially underwent TBCLB followed by video-assisted thoracoscopy for SLB at the same anatomical location. It was concluded that poor concordance could have been resulted in different management strategy in 11 of 21 (52%) of cases (50).

**Airway Management and Anesthesia**

There is wide variation in terms of where and how TBCLB is performed. Earlier studies reported airway management with rigid bronchoscopy under general anesthesia in case of major bleeding however safety of the procedure has been established in varies airways (Laryngeal mask anesthesia, endotracheal tube with conventional ventilation or Jet ventilation) and sedation level (general and conscious). Centers usually select the location, anesthesia type and airway modality based on their experiences. Our group has compared diagnostic yield and complication rate of TBCLB in ILD patients in a retrospective study and reported no difference in biopsy quality and diagnostic yield but increased trend of more bleeding requiring cold saline and topical epinephrine instillation (moderate bleeding) in endoscopy unit (30).

**Future Directions**

TBCLB is a safe and promising technique in the diagnosis of interstitial lung diseases. Variations in reporting and procedure itself makes difficult to compare published studies. Standardization in patient selection, indications, diagnostic yield, safety precautions and technical aspect will provide more robust information about its utility in diagnosis and other endpoints such as complication rates. Future comparative studies with surgical lung biopsy and unified data collection are particularly needed.

**CONFLICT of INTEREST**

The authors reported no conflict of interest related to this articles.

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