Robotic bronchoscopy for pulmonary lesions: a review of existing technologies and clinical data

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Abstract: Bronchoscopic interventions are preferred for sampling suspicious pulmonary lesions as they have lower complications and can achieve diagnosis and staging in one single procedure. Limitations in existing guided bronchoscopy platforms has led to developments in robotic assisted technologies. These devices may allow the bronchoscopist to more precisely maneuver the scope and instruments into the periphery of the lungs under direct visualization while also ensuring stability during sampling of the target lesions. These devices have the potential to improve the diagnostic yield in sampling peripheral lung lesions and may play a role in the treatment of non-operable or oligometastatic peripheral tumors using bronchoscopic ablative therapies. In this article, we review the existing robotic bronchoscopy technologies and summarize the available pre-clinical and clinical data supporting their use.

Keywords: Robotic bronchoscopy; peripheral lung lesion; lung cancer; biopsy; navigational bronchoscopy

Introduction

The detection of lung nodules has increased in the last decade owing to the increasing use of computed tomography (CT) and institution of the lung cancer screening programs. An estimated 1.6 million new pulmonary nodules are predicted to be detected every year in the USA (1). These statistics present a new challenge for physicians caring for these patients. The National Lung Screening and NELSON trials both showed that a significant number of patients had a positive screen, and majority of these nodules (~80%) were located in the periphery of the lung (2-4). The American College of Chest Physicians and National Comprehensive Cancer Network guidelines recommend diagnosis of the primary lesion, staging and obtaining tissue for genomic alterations and PD-L1 status using the least invasive modality and ideally in a single procedure (5,6). In many patients, therefore, an initial bronchoscopic approach facilitates both biopsies of the primary lesion as well as mediastinal staging prior to curative-intent therapy.

Rationale for developing new bronchoscopy platforms for peripheral lung lesions

The need to safely and effectively sample lung lesions, has led to the development of virtual bronchoscopy (VB), radial endobronchial ultrasound (r-EBUS) and electromagnetic navigation (EMN). The diagnostic yield of bronchoscopy using EMN ranges from 67–84%, with one study showing a diagnostic yield of 73% after one year of follow up (7). Regardless of the guided bronchoscopy platform, the diagnostic yield of bronchoscopy for lung lesions has generally remained lower than that of image guided transthoracic needle aspiration (92.1%) (8).
The lower diagnostic yield with conventional bronchoscopy could be explained by “getting lost” in the peripheral airways (4). EMN was thus introduced to guide navigation to these peripheral pulmonary lesions (PPLs) but lacked direct visualization of the airways. The use of EMN has been combined with r-EBUS to increase diagnostic yield but continues to be limited by respiratory motion and CT-to-body divergence. These limitations in guided bronchoscopy led to the introduction of the robotic assisted bronchoscopy (RAB) systems that allows the operator to navigate through smaller airways under direct visualization while continuing to offer either EMN guidance (Monarch™ platform by Auris Health Inc.) or Shape Sensing Technology (Ion™ Endoluminal System by Intuitive Surgical) to find target airways and provide stability during sampling of the target lesion.

Robotic bronchoscopy: a historical perspective

The first “robot surgeon” used on human patient was the PUMA 560 robotic system introduced in 1985 to perform CT guided neurosurgical biopsies (9). In the 1990’s, the AESOP (Automated Endoscopic System for Optimal Positioning) robotic platform became the first system approved by the FDA for its use during endoscopic surgical procedures. In 2000, the Da Vinci Surgery System was approved by the FDA for general laparoscopic surgery (10). Since then, the use robotics has expanded from the field of general surgery, to gynecologic surgery and urological surgery as well as cardiac and thoracic surgery and has become an integral part of surgical procedures in the United States. The first robotic system to be introduced in the field of bronchoscopy was the Monarch™ platform by Auris Health and received FDA approval in March 2018. Subsequently another robotic bronchoscopy platform, Ion™ Endoluminal System developed by Intuitive Surgical received FDA approval in February 2019. The introduction of robotic bronchoscopy to the field of Interventional Pulmonology has been a source of great excitement in the era of lung cancer screening.

Monarch system: summary of evidence from cadaver and patient studies

The Monarch™ platform by Auris Health includes an outer sheath and an inner bronchoscope with a 4-way steering control, electromagnetic navigation guidance and continuous peripheral visualization during navigation as well as biopsy (Figure 1). The first study of the Monarch™ platform was performed on cadavers to assess the reach of the RAB platform compared to a conventional thin bronchoscope with a similar outer diameter (4.2 mm). A guidewire was inserted into each segmental bronchus till it reached the pleura as noted on fluoroscopy. The conventional thin bronchoscope was then advanced over the guidewire as far as possible. The bronchoscope was then removed and the Monarch™ RAB system was advanced over the guidewire to its maximum extent. In each of the segments, the Monarch™ RAB platform had farther access to the periphery of the lung compared to the bronchoscope (9th vs. 6th airway generations). This was likely due to the structural support provided by the RAB system while the inner scope was advanced. In comparison, the conventional bronchoscope was noted to prolapse in the trachea when trying to be advanced to the apical segments. Another reason was likely the ability of the RAB system to negotiate subtle turns and bifurcations in the distal airways allowing for further maneuverability (11).

Rojas-Solano et al performed the first feasibility study using the Monarch™ RAB platform in 15 patients with suspicious central lesions or peripheral lesions with a bronchus sign. The average lesion size was noted to be 26 mm (10–63 mm) with 12 patients with peripheral lesions and 3 patients with a central lesion. R-EBUS was not used to assist sampling. Samples were obtained in 14/15 (93%) patients and none of the patients had a pneumothorax or significant bleeding as a complication (12).

The recent ACCESS trial studied the Monarch™ Auris RAB platform in 8 cadavers with artificial tumor targets created using the aqueous solution of a mixture of agar, gelatin, iodinated contrast and colored mica powder which were injected transthoracically into the lung parenchyma using fluoroscopic guidance. The mean nodule size was 20.4 mm (9.6–28.3 mm) with 45/67 (68%) nodules measuring 10–20 mm in diameter. The overall diagnosed yield was noted to be 94% (63/67) using transbronchial needle aspiration (TBNA) which was increased to 97% (65/67) by the use of transbronchial forceps (TBBX) (13). It was interesting to note that the yield for lesions with eccentric ultrasound (r-EBUS) view in this study was 97% (47/48) which is significantly higher than that is previously published in literature (30–40%) (14). This improvement may be explained by the fact that the RAB systems provide the ability to maneuver and control the distal end of the scope and provide the ability to aim biopsy instruments with greater precision through a uniquely designed catheter and...
A recent retrospective post-marketing multicenter study using the Monarch™ Auris robotic platform in 165 patients with 167 lesions showed successful navigation (acquisition of a r-EBUS view or diagnostic tissue on pathology) to 88.6% of the lung nodules (70.7% were located in the outer third of the lung) (16). The average lesion size was 25 mm (±15 mm) with 71% of the nodules under 30 mm and 63.5% of the lesions had a bronchus sign. The conservative diagnostic yield was 69.1% (patients with presence of inflammation on biopsy without any follow-up excluded) with a maximum diagnostic yield estimated to be 77%. Consistent with the results of the ACCESS trial, the diagnostic yield even with an eccentric view on r-EBUS was noted to be 71.1%, which is significantly higher than that previously described in the literature reporting on the various r-EBUS image patterns and diagnostic yields (14,17). The diagnostic yield of 54% was noted in the absence of the bronchus sign, which is also higher than the previously reported yield of 31–44% (18,19) but lower than that of the NAVIGATE study (7). A 3.6% rate of pneumothorax and 2.4% risk bleeding rate were comparable to other guided bronchoscopy studies. A major limitation of the retrospective series was that follow-up was only available for 6 months as compared to a 12-month follow up noted in major prospective trials.

A prospective single-arm multicenter post-marketing study called BENEFIT is currently assessing the performance of the Monarch™ Auris RAB platform in human subjects of biopsying peripheral lung lesions (https://clinicaltrials.gov/ct2/show/NCT03727425). The primary outcome measures include successful navigation to target pulmonary lesions and to assess device and procedure related adverse events. The preliminary results from this study were recently presented at the American College of Chest Physicians annual meeting and showed lesion localization in 96% cases in 56 patients enrolled in the
A multicenter prospective post marketing study (TARGET) is currently enrolling patients and aims to enroll 1,200 patients across 30 sites undergoing robotic assisted transbronchial lung biopsy over a period of 4 years. The primary endpoint includes device or procedure related complications, with secondary endpoints assessing the diagnostic yield with the patients being followed up to 24 months post-procedure (https://clinicaltrials.gov/ct2/show/NCT04182815).

**Ion system: summary of evidence from cadaver and patient studies**

The Ion™ Endoluminal System by Intuitive Surgical includes an articulating, flexible catheter with shape sensing technology, which provides positional and shape feedback along with a video probe for live visualization while driving the catheter (Figure 2).

Fielding et al recently published the first human study using the Ion™ Endoluminal System by Intuitive that included 29 patients with the mean lesion size of 12.2 mm (±4.2) in the axial plane with majority (68.9%) of the lesions being located in the upper lobes. Only 17 patients (58.6%) had a bronchus sign visible on CT with approximately half the cases having an eccentric r-EBUS view. The primary end point of reaching the target and obtaining samples was achieved in 28/29 (96.6%) of the cases. The overall diagnostic yield was 79.3% with a diagnostic yield for malignancy of 88% (21). The high yield, especially for cases with an eccentric r-EBUS view was attributed to the ability to visualize peripheral airways and the ability to deploy the TBNA needle perpendicular to the airway/bronchus, towards the lesion.

The cadaver-based Precision-1 study compared the ultra-thin bronchoscope using radial EBUS (UTB-rEBUS), EMN and RAB platforms (Ion™ Endoluminal System by Intuitive) in localizing and puncturing of artificially implanted small (<2 cm) peripheral nodules in cadavers. Majority (80%) of the nodules were placed in the periphery with the mean size of 16.5 mm (±1.5 mm) and 50% of the nodules had a positive bronchus sign. Nodule localization was achieved in 100% cases with the RAB platform vs 85% with EMN guidance and 65% with the use of UTB-rEBUS. Once the biopsy was performed using a needle, the relationship of the needle to the lesion was studied using cone beam CT. Successful puncture of the nodule (defined as the needle in the nodule or going through the nodule) was noted in 80% cases with the Ion™ RAB platform compared to 45% with EMN (P=0.022) and 25% with the use of UTB-rEBUS (P<0.001). Even in the lesions that were missed, the median needle to miss distance was noted to be 4 mm (3–5 mm) in the RAB platform compared to 7 mm (4–9 mm) with EMN guidance and 13 mm (6–28 mm) with the use ofUTB-rEBUS. The authors stated that the Ion™ RAB platform increased the ability to localize and precisely puncture peripheral nodules (22). The study has several limitations. There is no evidence to suggest that implanted lesions mimic real lung tumors, therefore the generalizability of this study’s findings is highly questionable. In addition, there are concerns about the validity of the EMN testing, as it appears that industry’s standard operating procedures (in regards to room mapping, etc.) were not followed in this study.

A prospective single-arm multicenter post-marketing study (PRECiSE) assessing the Ion™ RAB platform is currently in progress (https://clinicaltrials.gov/ct2/show/NCT03893539). This study aims at enrolling 360 patients with the primary outcomes assessing navigation and biopsy success and secondary outcomes assessing the associated complications with the use of this system.

![Figure 2 Ion™ endoluminal system by intuitive surgical. (A) Computed tomography (CT) image of the right upper lobe nodule (arrow); (B) target view, concentric radial EBUS view and fluoroscopic image of the robotic bronchoscope.](https://clinicaltrials.gov/ct2/show/NCT04182815)
Robotic-assisted bronchoscopy platforms: differences and similarities

Due to their stability, adjustable angulation and peripheral visualization, the robotic platforms have the ability to potentially overcome some limitations of the currently available guided bronchoscopy systems and increase the diagnostic yield. With one platform, there is increased structural support by the outer sheath that can be locked in the target segments (usually 3rd or 4th generation) before advancing the scope into the target airway. The scope itself also has an increased ability to make subtle turns due to the four way steering capability and has been proven to have a farther reach than the conventional bronchoscopy platforms in cadaver studies (11). The scope can be locked in position, while the instruments are advanced thus minimizing airway distortion. The platforms also allow for direct visualization of the peripheral airways, and with one system, direct visualization of the biopsy tools as they are advanced outside the working channel. This can ensure that the bronchoscopist continues to remain in the target airway and also enables the bronchoscopist to steer the tools towards the target (Figure 1). This is relevant for practice as studies to date have shown that the ability to maneuver the scope and precisely guide the instruments can help increase the yield in targets with an eccentric r-EBUS view. The small OD of bronchoscope also helps the scope to be wedged and locked in the target segment, which can allow for tamponade and containment of the bleeding. The published evidence, system specifications, accessory tools as well as the potential advantages and disadvantages of both RAB systems currently available commercially are summarized in Table 1.

Table 1 Comparison of the Monarch™ Platform and Ion™ Endoluminal System

| Robotic-assisted bronchoscopy platforms | The Monarch™ platform Auris Health Inc. | Ion™ Endoluminal System Intuitive Surgical |
|----------------------------------------|----------------------------------------|--------------------------------------------|
| **FDA approval** | March, 2018 | February, 2019 |
| **Studies** | • Chen et al. 2018 (REACH Cadaver Study) (11) | • Fielding et al. 2019 (Human Feasibility Study) (21); |
| | • Rojas-Solano et al. 2018 (Human Feasibility Study) (12) | • Yarmus et al. 2019 (Cadaver Study Comparing RAB vs. EMN vs. Conventional bronchoscopy with r-EBUS) (22); |
| | • Chen et al. 2019 (ACCESS Cadaver Study) (15) | • Ongoing—Multicenter prospective BENEFIT trial (https://clinicaltrials.gov/ct2/show/NCT03727425) |
| | • Chaddha et al. 2019 (Retrospective Post Marketing Study) (16) | • Ongoing—Multicenter prospective PRECIsE Trial (https://clinicaltrials.gov/ct2/show/NCT03893539) |
| | • Ongoing—multicenter prospective BENEFIT trial (https://clinicaltrials.gov/ct2/show/NCT03727425) | |
| | • Ongoing—multicenter prospective TARGET trial (https://clinicaltrials.gov/ct2/show/NCT04182815) | |
| **Bronchoscope specifications** | • Inner bronchoscope (4.2 mm) & Outer sheath (6 mm), both with 4-way steering control; | • 3.5 mm outer diameter fully articulating catheter; |
| | • The sheath can be locked in place and the bronchoscope can be advanced to the airways under EMN guidance and direct visualization; | • 2 mm working channel; the catheter has a shape-sensing fiber along its entire length which provides positional and shape feedback; |
| | • 2.1 mm working channel; | • Catheter articulates 180 degrees in any direction; |
| | • Constant peripheral visualization during workflow at the target | • Integrated vision probe that provides navigation; |
| | | • Vision probe has to be removed prior to tissue sampling |
| **Navigation technology** | Relies on Electromagnetic Navigation along with peripheral vision and real time input from the micro-camera at the tip of the bronchoscope | Relies on fiber optic sensing technology “shape sensing” and peripheral vision for navigation; |
| | | • The shape sensing technology is reportedly not sensitive to metal objects |
| **Instruments** | • Auris needle (currently not available on the market); | • Flexible needle—Flexision™; |
| | • Other needles such as Olympus Periview Flex or Arc Point SuperDimension needles; | • The biopsy needle can be visualized along its length, and its length can be set to avoid the pleura and reach the middle of the nodule; |
| | • Can use R-EBUS, needle, biopsy forces or brush through the working channel; | • Can use R-EBUS, needle, biopsy forces or brush; |
| | • The direction and positioning of the R-EBUS, needles, brushes or forces instruments can be re-oriented under direct guidance | relies on the fiber optic sensing technology as well as real time positioning during tissue sampling |

Table 1 (continued)
In addition to their potential for improving diagnostic yield when sampling peripheral lung lesions, the RAB platforms may guide ablative therapies for treating oligometastatic lesions or inoperable peripheral lung tumors. Both animal studies and case studies in humans have already demonstrated the feasibility of guided bronchoscopic ablative therapies such as photodynamic therapy (23), microwave ablation (24), radiofrequency ablation (25), laser interstitial thermal therapy (26), cryoablation, and bronchoscopic thermal vapor ablation. RAB platforms may increase the accuracy in guiding the catheters to peripheral lesions and provide a stable platform to deliver ablative therapies (4,27). It is possible that these interventions will be performed by combining the use of RAB platforms and direct imaging guidance such as cone-beam CT for precise application of the probes and for monitoring intra-procedural effect. The feasibility of combining robotic and cone beam CT technologies in routine clinical practice remains to be determined.

**Conclusions**

Based on the results of published evidence and our own experience, we believe that robotic bronchoscopy platforms are a step forward towards improving the diagnostic yield in sampling peripheral lung lesions. They also may have a potential role in the treatment of non-operable or oligometastatic peripheral tumors using bronchoscopic ablative therapy.

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