

# Transbronchial microwave ablation of lung nodules with electromagnetic navigation bronchoscopy guidance—a novel technique and initial experience with 30 cases

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**Background:** Microwave ablation of lung nodules may provide a faster, larger and more predictable ablation zone than other energy sources, while bronchoscopic transbronchial ablation has theoretical advantage of fewer pleural-based complications than percutaneous approach. Our study aims to determine whether the novel combination of bronchoscopic approach and microwave ablation in management of lung nodules is technically feasible, safe and effective.

**Methods:** This is a retrospective analysis of a single center experience in electromagnetic navigation bronchoscopy microwave ablation in hybrid operating room. Patients had high surgical risks while lung nodules were either proven malignant or radiologically suspicious. Primary endpoints include technical feasibility and safety.

**Results:** Total of 30 lung nodules from 25 patients were treated. Mean nodule size was 15.1 mm, and bronchus directly leads to the nodules (bronchus sign positive) in only half of them. Technical success rate was 100%, although some nodules required double ablation for adequate coverage. Mean minimal ablation margin was 5.51 mm. The mean actual ablation zone volume was -21.4% compared to predicted, likely due to significant tissue contraction ranging from 0-43%. There was no significant heat sink effect. Mean hospital stay was 1.73 days, and only 1 patient stayed for more than 3 days. Complications included pain (13.3%), pneumothorax requiring drainage (6.67%), post-ablation reaction (6.67%), pleural effusion (3.33%) and hemoptysis (3.33%). After median follow up of 12 months, none of the nodules had evidence of progression.

**Conclusions:** Bronchoscopic transbronchial microwave ablation is safe and feasible for treatment of malignant lung nodules. Prospective study on clinical application of this novel technique is warranted.

**Keywords:** Microwave ablation; transbronchial ablation; lung cancer; electromagnetic navigation bronchoscopy; hybrid operating room

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#### Introduction

With strong evidence for low-dose computer tomography (CT) scans screening in high risk populations (1), incidental discovery of lung nodule is becoming more common. Many of them are small, sub-solid, and harbour pre-malignant or early stage tumours. Local therapies as definitive treatment of these lesions is among the standard managements, especially in patients with surgical contraindications, for instance prohibitive cardiorespiratory risks or inadequate lung function. Among patients newly diagnosed with early stage lung cancer, 5-10% have absolute or relative contraindications to surgery (2,3), but the percentage could be as high as 28% in octogenarians (4). Treatment modalities such as stereotactic body radiation therapy (SBRT) (5), percutaneous ablation techniques for instance radiofrequency (RFA) (6), microwave (MWA) (7) and cryotherapy (8) are highly effective and are associated with reasonable local control rates ranging from 64% to 69.8% at 2 years. However, these procedures are also associated with complications: SBRT carries up to 22.3% risk of radiation pneumonitis and pneumonia (9), while percutaneous ablation techniques carry 11-52% risk of pneumothorax and bronchopleural fistula (BPF) (10). In terms of energy source, radiofrequency relies heavily on electrical conductance of tissues, thus the high impedance of human lung tissue may limit its effectiveness (11).

Bronchoscopic transbronchial ablation techniques utilizing radiofrequency energy for eradication of tumor cell have been described (12), with theoretical advantages of less pneumothorax/BPF due to non-trans-pleural route of entry, being able to reach peri-bronchial tissues easily, and having access to particular locations in lung which are otherwise difficult or impossible to reach. Meanwhile, microwave energy produces a larger and more predictable ablation zone, as it is minimally affected by impedance and has less heat-sink effects (11). Building upon our institute's experience in lung nodule dye-marking and biopsy via electromagnetic navigation bronchoscopy (ENB) (13,14), we identified a selected group of cases which are suitable for bronchoscopic transbronchial microwave ablation under ENB guidance. Being one of the first centres to perform ENB microwave ablation in clinical setting, we retrospectively analyze data from the first cohort of patients who had completed the procedure. Primary objective of this study is to assess the safety, feasibility and efficacy in reference to ablative effects and treatment outcomes. We present the following article in accordance with the STROBE reporting checklist (available at http://dx.doi.

org/10.21037/tlcr-20-1231).

# Methods

### Trial design

This study is a single-centre retrospective analysis of patients who underwent bronchoscopic transbronchial microwave ablation of malignant or suspicious lung nodules. The study was conducted in compliance with the Declaration of Helsinki (as revised in 2013) and approved by local institutional review board (the Joint Chinese University of Hong Kong – New Territories East Cluster Clinical Research Ethics Committee), CREC reference number of 2020.524. All cases were discussed in multi-disciplinary meetings with oncologists' input, and all study participants gave informed consent before taking part in the study.

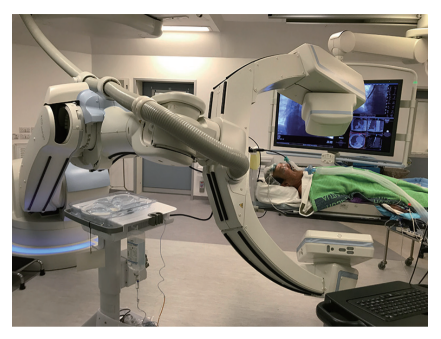
#### Enrolment criteria

Patients with radiologically suspicious lung nodules on CT scans and carried high surgical risk or have refused surgery were eligible for evaluation for ENB microwave ablation. Favourable radiologic features for selection included presence of bronchus sign (presence of bronchus directly leading to nodule), peripheral location, tumor size less than 3 cm, and at least 5 mm away from large blood vessels or mediastinal structures to avoid significant heat sink effect or thermal injury to other visceral organs. Special attention should be paid to avoid including brachial plexus or phrenic nerve into the designed ablation zone. Patients with biopsyproven lung cancer must have CT or PET/CT scans for exclusion of distant metastases, while nodal involvements were ruled out with endobronchial ultrasound biopsy if lymph nodes were large or hypermetabolic, such that eligible nodules are effectively T1N0M0. Ablation was offered to biopsy-proven lung metastasis if the primary tumour was treated with curative intent and all oligometastases in body could be treated. For the nodules which did not have histology either due to failed or refused biopsy, ablation was only offered if they were radiologically suspicious on serial imaging, and with patients' prior informed consent. During the period from March 2019 to June 2020, 25 patients with 30 nodules were included in the study.

#### ENB microwave ablation procedure

All patients had a plain fine-cut CT scan within 3 months from the scheduled ablation date for pre-operative planning

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**Figure 1** Set-up in the hybrid theatre is shown. The microwave catheter is connected to the microwave ablation console at the lower left in a particular direction such the C-arm was still able to rotate around the table to perform CBCT.

of ENB pathway.

Treatment procedure is summarized in Table S1. On the day of procedure, the patient was intubated inside our hybrid theatre with cone-beam CT (CBCT) and fluoroscopy capabilities. After the first CBCT, navigation to the target lung lesions was performed under ENB with SuperDimension<sup>TM</sup> Navigation System (Covidien<sup>TM</sup>, Plymouth, MN, USA) as per previously described (15) (Figure S1). Upon reaching segmental bronchus, a locatable guide (LG) within an extended working channel (EWC) was advanced through the bronchoscope following the navigational route until reaching the target (Figure S2). In cases where bronchus sign was absent and transbronchial access was required, CrossCountry<sup>TM</sup> Transbronchial Access Tool (Covidien<sup>TM</sup>, Plymouth, MN, USA) was utilized (Video 1). A needle was deployed into the lesion and a second CBCT was done to confirm desired location of needle tip within or closely abutting lesion. The CrossCountry<sup>TM</sup> dilator was then used to create the transbronchial access (Video 2). Biopsy was performed at this stage if necessary.

The needle and dilator were then exchanged to the Emprint<sup>TM</sup> Ablation Catheter with Thermosphere<sup>TM</sup> technology (Covidien<sup>TM</sup>, Plymouth, MN, USA) (Figure S3). The EWC was retracted to unsheathe the tip of microwave catheter (*Videos 3,4*), and a third CBCT was performed to

confirm accurate positioning of the catheter. An appropriate ablation energy was determined and predicted margin was measured. Microwave ablation was then carried out (*Figure 1*, Figure S3). After cooling for 10 minutes, the first post-ablation CT scan was done to assess the inclusion of lesion into ablation zone, the size of ablation zone and the minimal margin achieved. Double ablation may be performed by either re-ablating in same position, pull-back and re-ablate, or re-navigation and re-ablate (termed as "bracket ablation") (Figure S4).

#### Follow up, response evaluation and data analysis

Chest X-rays were performed on post-operative day 0 and 1, and patients were discharged earliest at 1 day after ablation if no complications arose. Post-ablation fine-cut plain CT scans were arranged at first month, 3<sup>rd</sup> month, 6<sup>th</sup> month, 9<sup>th</sup> month and 12<sup>th</sup> month, but the full set was not possible for some cases due to patient default or resource restrictions in our public hospital centre. Additional CT scans were done in case of any significant complications. Private scans including PET/CT or CT thorax with contrast were also evaluated if they were available for review. The response to ablation was evaluated by qualified radiologists for local control using the modified Response Evaluation

Criteria in Solid Tumours (mRECIST). Statistical analyses were performed with Microsoft Excel 2010 and IBM<sup>®</sup> SPSS<sup>®</sup> Statistics 20. Sample size was not estimated due to the retrospective nature of study. Patient and nodule characteristics were summarized with descriptive statistics, while post-ablation CT appearances and clinical outcomes were compared using Student's t test and chi-square test as appropriate. A P value of  $\leq 0.05$  was considered significant.

## Results

#### **Baseline characteristics**

Total of 25 patients were enrolled between March 2019 and June 2020. Patient characteristics are summarized in *Table 1* and Table S2. Thirty lung nodules were ablated, in which 5 patients had more than one lung nodule ablated. Concomitant ENB biopsy of target lesion was performed in 6 cases.

All patients had absolute or relative contraindications to surgery, including poor lung reserve or previous contralateral lobectomy, or comorbidities limiting survival, for instance significant cardiopulmonary disease or know metastatic disease. Fourteen patients (56%) had previous lung operations, 78.6% of those had previous contralateral lobectomies making single-lung ventilation potentially problematic. Patients included in the study had significant multiple comorbidities and the median Charlson comorbidity index was 6.

The mean maximal diameter of lung nodule size was 15.1 mm (range, 7 to 29 mm). Biopsy of lung nodules were either performed pre-operatively using CT-guided percutaneous biopsy, or during ENB procedure just before ablation. Pre-operative biopsy was not performed for 10 lung nodules mostly due to difficult access or patient refusal of biopsy. The histological types included lung adenocarcinomas (13.3%), atypia/pre-malignant (33.3%) and metastatic cancer (3.33%). Total of 15 nodules (50%) were negative for malignancy or no biopsy done but still radiologically suspicious. The mean distance between nodule and pleura is only 7.46 mm. Total of 6 nodules were abutting the pleura and 14 nodules were within 5 mm from pleura, making pneumothorax a potential concern.

# Procedural characteristics

Procedural characteristics were summarized in *Table 2*. After excluding cases with concomitant biopsy, the mean

total procedural time (from intubation to extubation) was 126.3 minutes. Intra-operative CBCT scans were used to plan, guide and adjust catheter locations, and the median number of CBCT required is 7.

Four nodules (13.3%) required planned or unplanned double ablation to include the nodule in a satisfactory ablation zone with adequate margin. Mean procedure time for single ablation was 116 minutes while that for double ablation was 160 minutes. Concomitant biopsy also significantly lengthens procedural time to mean of 165 minutes.

#### Immediate post-ablation statistics

A minimum of 5mm margin was planned for each ablated nodule. The expected ablation zone border was drawn using PURE<sup>®</sup> platform (Siemens Heathineers, Germany) software and the minimum predicted margin was measured (*Figure 2*), which had a mean ( $\pm$ SD) of 6.63 ( $\pm$ 2.89) mm. The mean ( $\pm$ SD) minimal actual margin after ablation was 5.51 ( $\pm$ 2.48) mm (*Table 3*), and there was no significant difference between the predicted and actual minimal margin (P=0.086). Technical success rate, defined as inclusion of lesion into ablation zone with adequate margin, was 100%.

Tissue contractions occur after ablation, and should be taken into consideration when determining whether adequate margin had been achieved. The contraction percentage is not a constant, but rather is inversely proportional to the distance from the centre of ablation (Figure 3A). This is the so-called "black hole" theory, meaning that contraction is more pronounced when a reference point is closer to the centre of ablation, in analogy of a black hole where objects closer to the centre experience a stronger attraction force. Each patient had different contraction curves, with examples shown in Figure 3B. When all patient's data was pooled on a scatter plot (Figure 3C), there is a rough maximal contraction percentage for each distance, for example a maximum of 40% contraction for a 20 mm distance from centre. In general, the contraction percentage ranges from 0-43%.

The post-ablation zone volume was generally smaller than the predicted volume, with a mean of -21.4% (SD =38.9%), but this was without consideration of contraction. The actual ablation zones were as predicted (9 cases) or larger (5 cases) in 46.7% of cases. The cases with smallerthan-predicted ablation zone were analyzed against possible factors affecting the ablation zone size, including presence of COPD, solidity of lesion, and presence of  $\geq 3$  mm blood vessels close to nodules, but none show statistically

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Table 1 Summary of patients and lung nodules characteristics

Table 1 Summary of patients and		Counts (%)	Mean	Range
Patients characteristics (25 pat	ients)			
Age (years)			68.8	51–86
Sex	Male	12 (48%)		
	Female	13 (52%)		
COPD		6 (24%)		
Smoking status	Current smoker	1 (4%)		
	Ex-smoker	6 (24%)		
	Never-smoker	17 (68%)		
Lung function	FVC		98.1% predicted	
	FEV1		94.4% predicted	
	FEV1/FVC ratio		0.732	
	DLCO		81.3% predicted	
Charlson comorbidity index			5.73	
Reason for ablation	Previous major lung resection	14 (56%)		
	History of non-pulmonary cancers	9 (36%)		
	History of major operation	22 (88%)		
	COPD stage 2 or above	4 (16%)		
	Expected limited survival / known metastatic disease	5 (20%)		
	Frailty/age >80 years	4 (16%)		
Lung nodules characteristics (3	0 lung nodules)			
Maximum nodule size (mm)			15.1	7–29
Lobe	RUL	9 (30%)		
	RML	3 (10%)		
	RLL	4 (13.3%)		
	LUL	8 (26.7%)		
	LLL	6 (20%)		
Location in lung	Periphery	22 (73.3%)		
	Middle one-third	8 (26.7%)		
	Innermost one-third	0		
Histology	Lung cancer	4 (13.3%)		
	Proven metastasis	1 (3.33%)		
	Atypia/pre-malignant	10 (33.3%)		
	Negative for malignancy	5 (16.7%)		
	Not available	10 (33.3%)		

Table 1 (continued)

Table 1 (continued)

		Counts (%)	Mean	Range
Suzuki class	Class 1–4	17 (56.7%)		
	Class 5–6 (predominantly solid)	12 (40%)	12 (40%)	
	Cavitatory	1 (3.33%)		
Bronchus sign	positive	15 (50%)		
	negative	15 (50%)		
Distance to fissure (mm)	ance to fissure (mm)		27.8	0–58.5
Distance to pleura (mm)			7.46	0–27.4
Blood vessel (diameter >3 mm)	Inside nodule	6 (20%)		
	Within 5 mm of nodule	12 (40%)		

LUL, left upper lobe; RUL, right upper lobe; RML, right middle lobe; LLL, left lower lobe; RLL, right lower lobe. COPD, chronic obstruction pulmonary disease. FVC, forced vital capacity; FEV1 forced expiratory volume in 1 second; DLCO, diffusing capacity for carbon monoxide

Table 2 Complications from bronchoscopic microwave ablation

Complications	Counts (per-lesion basis)	CTCAE grade	
Pain	4 (13.3%)	1	
Pneumothorax <sup>1,2</sup> requiring chest drain insertion	2 (6.67%)	3	
Post-ablation reaction/fever	2 (6.67%)	1	
Hemoptysis	1 (3.33%)	1	
Infected effusion	1 (3.33%)	1	
Abscess	0	_	
Bronchopleural fistula	0	-	
Hemothorax	0	-	
Arrhythmia	0	-	
Pneumonia	0	-	
Pulmonary edema	0	-	
Complications related to intubation and bronchoscopic manipulation (e.g., hoarseness, tracheobronchial injury)	0	-	
Unintended microwave burn (e.g., skin burn, vessel thrombosis, embolism, diaphragmatic paresis)	0	-	
Procedure-related mortality	0	_	

<sup>1</sup>Case 8 had intra-operative pneumothorax, required intra-operative chest drain insertion, which was removed on post-operative day 1 and discharged the same day. The peripheral lesion touched both the pleura and fissure, and review of CBCT showed inadvertent puncture of pleura during unsheathing (and unintentional advancement) of microwave ablation catheter. <sup>2</sup>Case 9 had pneumothorax on post operative day 1 and also required chest drain insertion. The initial lesion was cavitatory, and the ablation zone is much larger than expected (reaching pleura). He was later complicated by bacillus infection of pleural fluid and completed a course of intravenous antibiotics. Duration of stay was 16 days. CTCAE, common terminology criteria for adverse events.

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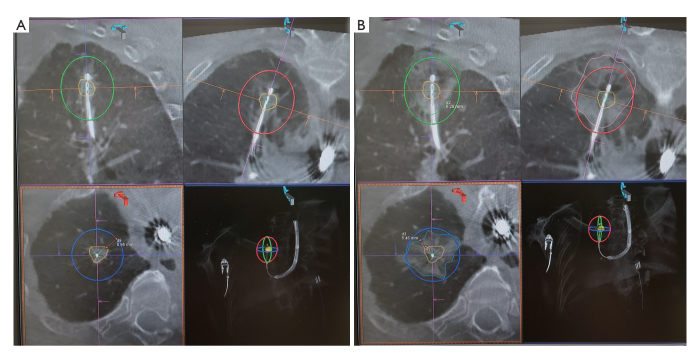


Figure 2 (A) This shows the target lung lesion (yellow tracing) in 3 axes before ablation. With ablation energy of  $100 \text{ W} \times 10$  minutes, the predicted ablation zone (red, green and blue ovals) is a spheroid measuring 42×35×35 mm, and the minimal predicted margin is 8.96 mm. (B) Figure is the 10-minute post-ablation scan, showing a minimal actual margin of 5.46 mm due to irregular shape of resultant ablation zone (red, green and blue irregular contour tracings), although the total ablation zone volume is similar to prediction.

significant results.

Regarding heat sink effect, there is no statistically significant correlation between a smaller ablation zone with the presence of a sizeable blood vessel (diameter  $\geq 3$  mm) within 5mm of nodule. Presence or absence of bronchus sign does not significantly affect fluoroscopy time, radiation dose, procedure time or the number of CBCT performed. In terms of learning curve, the first and second half of 22 ablation-only cases were compared. There was a trend towards shorter mean procedural time in the latter half of cases (139 vs. 111 minutes, P=0.075) (Figure S5).

#### Discharge statistics, safety and complications

The median length of hospital stay is 1 day, while the mean is 1.73 days. The great majority of cases (23 cases) were discharged on post-operative day 1, while 6 cases stayed for 2-3 days due to frailty or unrelated complaints.

There was a low rate of complications (Table 2). Mild immediate or delayed pleuritic chest pain was present in 4 cases (13.3%) and responded to common oral analgesics and did not prolong hospital stay. All 4 of the lesions were close to pleura, and the mechanism of pain could be due to pleural irritation after ablation of peripherally located lung nodules. No referred pain to shoulder due to phrenic nerve injury, or referred pain to arm due to brachial plexus injury occurred. Pneumothorax occurred in 2 cases (6.67%) and both required chest drain insertion. Post ablation reaction including fever occurred in 2 cases (6.67%); while mild hemoptysis was noted in 1 case (3.33%). There were only 2 CTCAE (common terminology criteria for adverse events) grade 3 events.

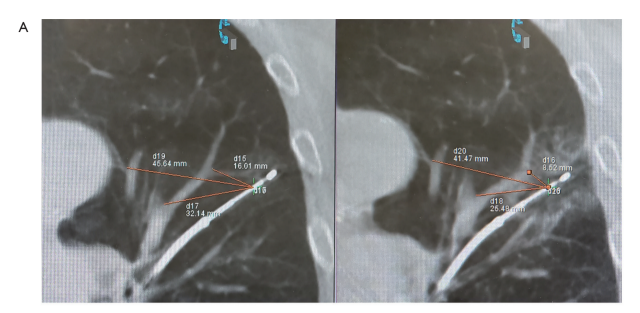
## Follow up and response

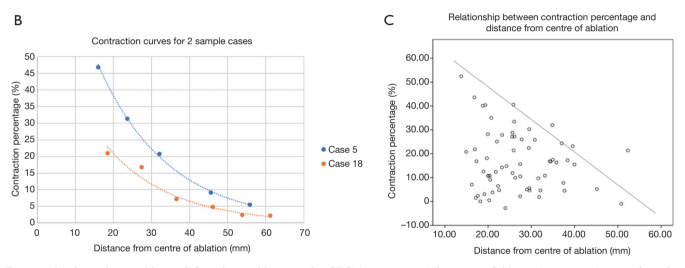
Median duration of follow up was 349 days (range from 132 to 615 days). Fifteen cases have completed at least 6-month post-ablation CT scans, with 8 of them already had post-ablation 1 year CT (Table 4). Complete response was achieved in 1 patient, partial response in 12 patients, and stable disease in the remaining 2 patients. There was no progressive disease identified. One unrelated death (case 2) occurred at 12 months after ablation. Morphology of ablated tumour on CT at one month post-ablation is

## Table 3 Ablation statistics

	Counts	Mean	Range
Single ablation	26		
Double ablation			
Re-ablate in same location	2		
Pull-back & ablate	1		
Re-navigate & ablate	1		
Number of ablations per lesion		1.13	1–2
Concomitant biopsy	6		
Adenocarcinoma	1		
Atypia	4		
Negative for malignancy	1		
Procedural time (minutes)			
All cases		134.5	70–200
Ablation-only		126.3	70–200
Single ablation		116.2	70–160
Number of CT			
All cases		7.6	4–12
Ablation-only		7.4	4–12
Single ablation		7.1	4–10
With biopsy		8.3	4–11
Catheter position			
Within centre of nodule	17		
Touching nodule border	10		
Outside nodule	3		
Ablation energy (J)		46,803	24,000–120,000
Duration of ablation (minute)		8.32	4–20
Power of ablation (Watt)		97.8	45–100
Radiation dose (µGym²), all cases		27,869	13,958–44,867
Margin (mm)			
Minimal predicted		6.63	3–14
Minimal actual <sup>1</sup>		5.51	3–10
Maximal actual		16.5	13–19
Actual AZ compared to predicted AZ			
Similar	9 (30%)		
Smaller	16 (53.3%)		
Larger	5 (16.7%)		

AZ, ablation zone. <sup>1</sup>Minimal actual margin is defined as the shortest distance between tumour border and ablation zone edge after 3-dimensional overlaying of ablated region on pre-ablation CT scan.





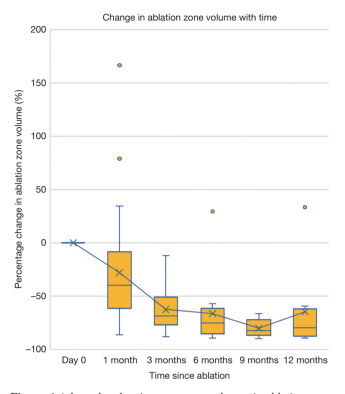
**Figure 3** (A) shows the pre-ablation (left) and post-ablation (right) CBCT appearances. The centre of ablation (green cross) is 1 cm from the tip of the microwave catheter as defined by the manufacturer. The distance between the centre of ablation and reference points (usually the bifurcation of recognizable neighbouring blood vessels) are measured pre- and post-ablation respectively. The contraction percentage for a particular distance from centre is calculated by (Pre-ablation distance – Post-ablation distance)/Pre-ablation distance ×100%. For example, contraction percentage is  $(32.14-25.48)/32.14\times100\% = 20.7\%$  for the point d17. (B) shows the contraction percentages measured at 5-6 reference points for two cases. References points are usually recognizable bifurcations of adjacent blood vessels which can be identified in both pre- and post-ablation CTs. The contraction percentage and distance from centre of ablation energy used, although the shape of curve should be similar. (C) shows a scatter plot of contraction percentage against distance from centre of ablation when data from all patients are pooled together. It is not a linear relationship, but there is a maximum possible contraction percentage for each distance. For instance, for reference points 20 mm from centre of ablation, a maximum of 40% contraction is observed. Likewise, for a reference point at 40 mm, a maximum of roughly 20% contraction is expected.

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Table 4 Short to	medium t	term follow	up results

Case	Percentage c	Percentage change (%) of AZ volume compared with immediate post-ablation CT scan				mprojor
	1m CT	3m CT	6m CT	9m CT	12m CT	- mRECIST
1	+78.9	-11.7	+29.4	_	+33.5	CR
2	-61.8	_	-85.4	-83.9	Death	PR
3	-81.1	-87.8	_	-	-82.7	PR
4	-	-63.8	-69.6	_	-	PR
5	-61.3	_	_	_	-89.1	PR
6	+34.6	-63.2	_	-66.0	-79.6	PR
7	-86.2	_	_	-89.5	-89.1	SD
8	-61.3	-73.2	-61.5	-	-69.5	SD
9	+166	-28.1	-79.6	-77.7	-59.3	PR
10	-16.4	-66.8	-74.9	-82.1	-	PR
11	-86.1	-	_	-	-	
12	-5.78	_	-68.2	_	-78.9	PR
13	-15.4	-69.6	_	-	-	
14	-71.2	-	-89.3	-	-	PR
15	-59.3	-77.0	-87.3	-	-	PR
16	-31.0	-30.1	-85.2	-	-	PR
17	-35.2	-	-56.8	-	-	PR
18	-72.9	-76.3	_	-	-	
19	-50.0	-84.0	_	-	-	
20	-3.85	-46.6	_	_	-	
21	-48.7	-74.8	_	-	-	
22	-53.1	-83.6	_	_	-	
23	+1.84	-	_	-	-	
24	-18.2	-	_	-	-	
25	-36.4	-	_	-	-	
26	-51.6	-	_	-	-	
27	-26.5	-62.3	_	-	-	
28	+17.1	-	_	-	-	
29	-	-	_	-	-	
30	-42.7	-	_	-	-	
Mean	-27.7	-62.4	-66.2	-79.8	-64.3	

AZ, ablation zone; mCT, months after ablation CT; mRECIST, modified response evaluation criteria in solid tumours; CR, complete response; PR, partial response; SD, stable disease.



**Figure 4** A box-plot showing percentage change in ablation zone from day 0 to 1 year after ablation. The rate of decrease in ablation zone volumes is initially fast and flattens out after 6 to 9 months, reaching a maximum of -79.8% at 9 months with our current data.

summarized in Table S3.

Regarding the ablation zone volumes in follow up CTs, there is a mean shrinkage of 27.8% when comparing first month CT to immediate post-ablation CT, however the variance is quite large. 60% of cases became smaller, 23.3% had static sizes. For the 10% of cases which got larger, on average there was a 93.4% increase in size, and were mostly those showing morphology of cavities. After that, the ablation zone continued to shrink, at a faster rate initially and slowing with time, eventually stabilizing in size by 9<sup>th</sup> to 12<sup>th</sup> month. In our series, compared to first month CT, ablation zone volume reduced by 66.2% at 6<sup>th</sup> month, and 64.3% by 12 months (*Figure 4*).

## Discussion

This retrospective study has demonstrated safety and feasibility of bronchoscopic transbronchial microwave ablation in management of lung nodules. Microwave energy for lung ablation demonstrates superior properties over its predecessor radiofrequency energy. Although both are electromagnetic energy that utilizes thermal heat to achieve tissue coagulation necrosis and cell death, radiofrequency energy induces frictional heating which occurs within only a few millimeters of the electrode, and the ablation zone size is mainly enlarged by thermal conductance, but charred and desiccated tissue surrounding the electrode increases electrical impedance (16). In contrast, microwave energy devices directly heat tissue to lethal temperatures through dielectric hysteresis (17). Being less dependent on electrical conductance, microwave energy deposition has less heatsink effect (17), is less susceptible to tissue impedance, and is able to produce faster, larger and more predictable ablation zones (11). In our series, the presence of nearby blood vessel did not significantly affect ablation zone volumes, although a mild local reduction in ablated area can sometimes be seen in certain cases.

Percutaneous microwave ablation for lung tumours produced reasonable results, with cancer-specific survivals ranging between 69–84.3% at 1 year, 42.1–61% at 3 years, recurrence rate of 44% at 3 years, while median time to local tumour recurrence was between 22.6–62 months (7,18-22). In a retrospective study comparing percutaneous microwave ablation and lobectomy for stage 1 non-small cell lung cancers (NSCLC), there was no significant difference in overall survival and disease-free survival (23).

Most of the thermal ablative techniques in literature applied the electrode in a percutaneous manner. Lately, transbronchial ablation techniques using different energy source have been researched in animal lung models (24-26). In humans, bronchoscopy-guided transbronchial ablation have been attempted by a Japanese and a Chinese group, utilizing RFA energy with reported safety (27-29). More recently, small case series of image-guided transbronchial microwave ablation of lung tumours have been reported by 2 separate groups with favourable benefit-risk profiles (30,31). Compared to percutaneous techniques, a major advantage of bronchoscopic ablation is avoidance of pleural puncture, and hence fewer pleural-based complications, including pneumothorax (6,10,32,33), bronchopleural fistulae (34) and needle tract pleural seeding (35). Another edge of bronchoscopic ablation is its ability to reach certain regions of lung which are otherwise difficult or dangerous for percutaneous route, for instance areas near mediastinal pleura or diaphragm, lung apex, and areas shielded by scapula. With evidence of safety and technical success of bronchoscopic microwave ablation in animal models (36),

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the authors' institute is one of the first to have performed ENB-guided microwave ablation on patients.

Our data supports the safety of bronchoscopic ablation compared to percutaneous route. Only 2 cases (6.67%) developed pneumothorax, and one of the pneumothorax may have been avoided if the catheter had not been advanced so far in as to puncture the pleura. The other pneumothorax likely resulted from rupture of a rapidlyenlarging cavity. The ablation zone reached pleura in 66.7% of our cases, indicating that an ablated pleura, as long as not punctured, is generally safe and does not produce pneumothorax. There was only 1 case (3.33%) of pleural effusion necessitating drainage, lower than the 6-19% reported in percutaneous ablation (10). The rate of other complications, 6.67% for post-ablation reaction, 3.33% for hemoptysis/bronchopulmonary haemorrhage, and 3.33% for infective complications, were comparable to percutaneous ablation (10).

Only selective cases are suitable for bronchoscopic transbronchial microwave ablation. Numerous lung nodule ablation studies have correlated risk of local recurrence with nodule size, with cut-off of 1.5, 2 and 3 cm respectively (6,7,19). It should be noted that the typical early postablative morphology of concentric GGOs on CT actually contains an outer rim of denser GGO containing congested lung tissue that retains viability (37). The size seen on CT scan overestimate the area of true coagulation necrosis by 4.1 mm (38) or even up to 8–9 mm (39). Therefore, multiple authors recommended a margin of at least 5 mm on CT scan assessment to ensure adequate tumour kill (18,38,40). For the Emprint microwave catheter used in the present study, the spheroidal maximal ablation zone width is 35 mm at 100 W for 10 minutes, being derived from in-vivo experiments in healthy porcine lungs (Figure S6). Taking into consideration a desired margin of at least 5mm on either side of lesion, the maximal nodule size suitable for ablation should be about 25mm, unless double ablation is planned.

The actual ablation zone volumes immediately postablation were generally smaller by a mean of -21.4%when compared to the predicted ablation zone dimensions provided by the manufacturer. A further analysis of the one-month ablation zone volume revealed a large variance in ablation zone sizes, ranging from -86.2% to +166%. The size of tumors, solidity of nodules and the presence of nearby blood vessels were not predictive of ablation zone sizes. The predicted ablation zone dimensions were derived from manufacturer's pre-clinical study in living healthy swine. The mismatch between predicted and actual ablation zone may be due to variations in lung properties between human and porcine lung (41), and differences in impedance and conductivity between normal lungs and tumour tissues.

It has been observed that tissue contraction occurred in most thermal ablation in different organs (16). To date only limited information on tissue contraction in human lung ablation is available. We found that contraction percentage is higher in the centre as opposed to periphery (black hole theory) (*Figure 3B*), but exact contraction percentage varies between individuals (*Figure 3C*). This may be explained by the differences in architecture and water content (16), lung nodule characteristics, and presence of surrounding blood vessels. Our findings may have significant implications on ablative margin and ultimately the technical success. By taking this phenomenon into account, clinicians may determine the ablative volume more accurately and reduce the risk of overcompensation with additional ablations.

Our study is limited by its retrospective nature, absence of parallel arm, and lack of histology for some lesions. Moreover, the contrasting effects of underestimation of ablation zone by CT due to tissue contraction and the overestimation of ablation zone by CT appearance indicates that CT is only fairly reliable in assessing and monitoring patient's response to ablation. Multi-phase contrast-enhanced CT, PET/CT and dynamic contrastenhanced MRI are better imaging modalities (37,42) but are not employed in the current study due to their limited availability. Nevertheless, current data is encouraging and support future prospective study on bronchoscopic microwave ablation. It will also be important to investigate the key parameters affecting ablation zone size such as lung densitometry and water content measurements. Long term results including local control and survival rates are also key endpoints to observe. Combined with simultaneous intratumoural chemotherapy (43), or enhanced permeability and retention (EPR) effect via hyperthermia induced by ablation (44), transbronchial lung cancer ablation will likely become an important part of the armamentarium in an exciting era of personalized cancer treatment.

## Conclusions

Bronchoscopic transbronchial microwave ablation is safe, feasible and potentially effective. This novel technique may represent a future treatment modality for early stage lung cancers and lung metastases. Together with ENB biopsy and intra-operative pathological evaluation, bronchoscopic microwave ablation in the hybrid theater is capable of providing a one-stop diagnosis and treatment for suspicious lung nodules (45).

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*Ethical Statement:* The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in compliance with the Declaration of Helsinki (as revised in 2013) and approved by local institutional review board (the Joint Chinese University of Hong Kong – New Territories East Cluster Clinical Research Ethics Committee), CREC reference number of 2020.524. All cases were discussed in multi-disciplinary meetings with oncologists' input, and all study participants gave informed consent before taking part in the study.

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