Airway Y-stent Insertion in Tracheobronchomalacia: A New Experience

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ABSTRACT:
This case report describes a rigid bronchoscopy-assisted placement of a silicone airway Y-stent in a patient who developed tracheobronchomalacia following repeated dilatations for post-tuberculous airway stenosis. This is the first report of an airway Y-stent insertion in Malaysia.

KEYWORDS: Tracheobronchomalacia, airway stent, tuberculosis.

INTRODUCTION

Tracheomalacia (TM) is a condition characterized by weakness of the trachea due to impaired supporting structures, i.e. the longitudinal elastic fibres of the pars membranacea and cartilaginous tissue, leading to a softer and more collapsible airway. If the bronchus is also involved, the term tracheobronchomalacia (TBM) is used. We describe a case of a lady, who had endobronchial tuberculosis complicated by tracheobronchial stenosis. She underwent repeated bronchoscopy-assisted dilatations using Controlled Radial Expansion (CRE) (Boston Scientific, Massachusetts, USA) balloon. Following repeated dilatations, she developed tracheobronchomalacia, which required rigid bronchoscopy-assisted insertion of an airway Y-stent.

CASE REPORT

The patient was a 38-year-old lady who presented with chronic cough and stridor. Her sputum acid-fast bacilli examination was negative. The initial flexible bronchoscopy found a lower tracheal stenosis of 4 cm in length, a stenosed and swollen right main bronchus, and a pin-hole stenosis of the right bronchus intermedius. She was diagnosed with endobronchial tuberculosis based on clinical and radiographic features. She received oral prednisolone for one month and anti-tuberculous therapy for a total of nine-month duration.

However, she was still stridorous and had reduced effort tolerance even after completion of treatment. A repeat bronchoscopy showed mild-to-moderate stenosis of the lower trachea measuring 2.5 cm in length and stenosis of the right bronchus intermedius, which was dilated using the Controlled Radial Expansion (CRE) balloon to 6 mm. Subsequently, she had seven sessions of CRE dilatations performed on the right bronchus intermedius, which was dilated to the maximum of 12 mm. She also had two CRE dilatations of the lower trachea to the maximum of 10 mm. Her symptoms improved after each dilatation procedure but always recurred a few months later. Just prior to the last CRE dilatation, her spirometry showed FEV1 0.38 L (14% of predicted), FVC 2.10 L (65% of predicted) and FEV1/FVC ratio of 18.1%. She could walk 180 m on the 6-minute walk test (6MWT) without dropping her oxygen saturation (pre-6MWT SpO2 98% and heart rate 79 beats/minute, post-6MWT SpO2 99% and heart rate 94 beats/minute).

In view of recurrent symptoms and airway stenosis, she was referred for airway stenting. Under general anaesthesia, a rigid bronchoscopic examination showed mild stenoses of the lower trachea (measuring 5.5 cm in length and 10 mm in diameter) and right bronchus intermedius. She also had severe lower tracheomalacia and right main bronchomalacia: The tracheal and bronchial endoluminal space was narrowed because opposing walls were partially collapsed causing partial occlusion of the airway (Figure 1).

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Figure 1: Tracheomalacia causing narrowing of tracheal lumen
Serial rigid endoscopic dilatations of the lower trachea were performed using rigid bronchoscope size 10 mm, 12 mm and 13.2 mm. Mild laceration and minimal bleeding were noted in the proximal end of the tracheal stenosis. A silicone airway Y-stent (Novatech, Boston Medical, Massachusetts, USA) size 15-12-12 mm was inserted and both bronchial sleeves were deployed satisfactorily (Figure 2). Right upper lobe bronchus was mildly oedematous due to stent manipulation. She was extubated immediately after the procedure and transferred to the ward for post-procedural recovery and observation.

A post-procedure chest X-ray revealed mild pneumomediastinum, which resolved spontaneously on the subsequent chest radiograph a week later. She was given I.V. dexamethasone 8 mg three times a day for the first three days and continued oral dexamethasone 8 mg three times a day for a total duration of one week. She also received I.V. ceftriaxone 1 g daily for one week. Spirometry performed a week after the procedure showed marked improvement: FEV1 1.99 L (69% of predicted), FVC 2.51 L (73% of predicted) and FEV1/FVC ratio of 79%. She was no longer stridorous and her functional status improved. She could attend to her usual daily activities, which she required assistance from a caregiver previously.

She was discharged well and planned for reassessment of her airway after six months. In the interim, she frequently complained of excessive secretions, which were alleviated using regular nebulized saline. Check bronchoscopy performed every one to two months also showed moderate build-up of granulation tissue at the lower end of the right limb of the airway Y-stent.

Six months later, she underwent repeat rigid bronchoscopy for reassessment of her airway. The airway Y-stent was seen in-situ. After removal of the airway Y-stent, significant lower tracheomalacia and right main bronchomalacia were still present. There was a moderate amount of granulation tissue in the right main bronchus and only minimal amount seen in the left main bronchus. Photocoagulation and grasp forceps were used to remove the excessive granulation tissue build-up in the right main bronchus. A new airway Y-stent (Novatech, Boston Medical, Massachusetts, USA) size 16-13-13 (cut to 5.8 cm (trachea) - 0.6 cm (left limb) - 1.0 cm (right limb) in lengths) was inserted. The patient recovered from the procedure and was discharged well.

DISCUSSION

TM and TBM in adults can be classified into primary (congenital) and secondary (acquired) forms.1,2 Congenital causes include polychondritis, idiopathic “giant trachea” or Mounier-Kuhn Syndrome while acquired causes include post-traumatic, e.g. post-intubation, post-traumatic and post-lung transplantation, chronic infection, chronic inflammation and chronic external compression of the trachea from aneurysms, tumours, cysts and abscesses.1

Although endoluminal tuberculosis leads to airway stenosis, it can also cause weakening and destruction of the airway and supporting cartilages. According to Feist et al.,2 the most common cause of acquired malacia of the airway is tracheostomy or intubation with endotracheal tubes. Possible mechanisms that were implicated in causing tracheal weakness include pressure necrosis, disruption of blood supply, friction on, and inflammation of, the mucosa.1

In this case, the patient’s tracheal and bronchial walls were weakened from the tuberculous disease process. With further repeated attempts at tracheobronchial dilatation, the stenosed segments were temporarily relieved but at the expense of worsening malacia of the affected tracheobronchial segments. This was seen on rigid bronchoscopic examination, which revealed only mild stenotic disease but severe lower TM and right main bronchomalacia. Severity of TM can be classified into mild, moderate or severe depending on the degree of tracheal obstruction during expiration.3 The TM is considered mild if the obstruction is one-half of the lumen, moderate if it is three-quarters and severe if the anterior and posterior walls touched.

Airway stents can be inserted to keep the malacic airway’s patent. Silicone stents are preferred to metallic ones because the latter are subjected to excessive granulation tissue formation and material fragmentation with time, which can cause airway obstruction, perforation and death.1 Furthermore, metallic stents are harder to remove and make future surgical interventions difficult.
The patient had a silicone Y-stent placed in the lower trachea and the sleeves deployed into the main bronchi. Our PubMed database search had shown no previous report of an airway Y-stent insertion under rigid bronchoscopy in Malaysia. Pang et al. described ten cases of self-expandable metallic stents inserted through flexible bronchoscopy but they were all straight airway stents.4

The patient’s symptoms and spirometry results showed significant improvement immediately following the stent placement. She will require long-term follow-up with bronchoscopy and review of her symptoms and lung function. One study reported that at mean 15 months after stent placement for stenotic and malacic lesions, peak flow parameters had declined though patients’ subjective improvement continued.5 In suitable candidates, surgery i.e. tracheoplasty, may also be considered as a definitive treatment option besides long-term stenting.1

REFERENCES
