CASE REPORT

Expiratory central airway collapse during positive pressure ventilation: a case report

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Abstract

Background: Physiologic narrowing of the central airway occurs during expiration. Conditions in which this narrowing becomes excessive are referred to as expiratory central airway collapse (ECAC). ECAC has a broad range of clinical presentations and is likely underdiagnosed [1]. It is usually managed with positive pressure ventilation, which acts as a pneumatic stent. Herein, we discuss the unusual clinical presentation of a patient with undiagnosed ECAC undergoing nonthoracic surgery under general anaesthesia.

Case presentation: We present the case of a 55-year-old man admitted for the placement of a ureteral JJ stent. Rapid desaturation occurred a few minutes after the onset of anaesthesia. After excluding the most common causes of desaturation, fibreoptic bronchoscopy was performed through the tracheal tube and revealed complete collapse of the left main bronchus. The collapse persisted despite the application of positive end-expiratory pressure and several recruitment manoeuvres. After recovery of spontaneous ventilation, the collapse was lifted, and saturation increased back to normal levels. No evidence of extrinsic compression was found on chest X-rays or computed tomography scans.

Conclusion: Cases of unknown expiratory central airway collapse reported in the literature were usually managed with positive pressure ventilation. This approach has been unsuccessful in the case described herein. Our hypothesis is that mechanical bending of the left main bronchus occurred due to loss of the patient’s natural position and thoracic muscle tone under general anaesthesia with neuromuscular blockade. When possible, spontaneous ventilation should be maintained in patients with known or suspected ECAC.

Keywords: Expiratory central airway collapse, Airway obstruction, Desaturation, Case report

Background

Physiologic narrowing of the central airway occurs during expiration. Conditions in which this narrowing becomes excessive are referred to as expiratory central airway collapse (ECAC). ECAC has a broad range of clinical presentations and is likely underdiagnosed [1]. It is usually managed with positive pressure ventilation, which acts as a pneumatic stent. Herein, we discuss the unusual clinical presentation of a patient with undiagnosed ECAC undergoing nonthoracic surgery under general anaesthesia.

Case presentation

A 55-year-old man (95 kg, 170 cm, BMI 32.9 kg.m⁻²) was admitted for emergency placement of a JJ stent. The patient had cerebral palsy, intellectual disability, deafness, and left hemiparesis due to prematurity and neonatal distress. He had acquired chronic obstructive pulmonary disease (COPD) with recurrent bronchial infections, pulmonary aspiration, and bronchospasms. He also had
sleep apnoea syndrome but was not adherent to con-
tinuous positive airway pressure (CPAP) treatment.

Cardiac auscultation was normal, and pulmonary aus-
cultation revealed clear, symmetrical vesicular murmurs,
with some added rhonchi.

The anaesthesia plan consisted of general anaes-
thesia in the supine position with orotracheal intubation.
After preoxygenation, the patient was given 7.5 μg of
sufentanil, 150 mg of propofol, and 35 mg of rocuronium
intravenously. Orotracheal intubation occurred without
incident, and auscultation showed bilateral and symmet-
rical pulmonary ventilation. SpO₂ was 100%. The venti-
lator was set to pressure controlled ventilation – volume
guaranteed mode with the following settings: respiratory
rate 12 min⁻¹, maximum pressure 30 mmHg, tidal vol-
ume 500 mL, positive end-expiratory pressure (PEEP)
5 cm H₂O. Anaesthesia was maintained with 1.5–2.0%
sevoflurane.

A few minutes after induction, haemoglobin oxygen
saturation rapidly dropped to 85%. Electrocardiogram
was normal with a heart rate of 90 beats per minute, non-
invasive blood pressure was 85/55 mmHg, FiO₂ was 45%,
end-tidal CO₂ at 33 mmHg with a normal square curve,
sevoflurane at a minimum alveolar concentration of 0.9.
There was no significant change in peak inspiratory pres-
sure. Based on these parameters, most causes of acute
desaturation were quickly ruled out.

Manual ventilation was resumed with FiO₂ 100%. On
re-auscultation, the absence of ventilation of the left
lung was observed. The first hypothesis was therefore
selective intubation of the right main bronchus. The
tube was withdrawn under direct laryngoscopic con-
trol until the tracheal cuff was positioned immediately
under the vocal cords. The integrity and pressure inside
the cuff were also checked.

Several recruitment manoeuvres (PEEP > 30 cm
H₂O for 40 s) were performed: SpO₂ transiently
increased to 94% before dropping again. Despite all
of these interventions, left lung ventilation remained
inaudible on auscultation, even during recruitment
manoeuvres. Fibreoptic bronchoscopy through the
tracheal tube revealed a complete collapse of the
left main bronchus (Fig. 1). Interestingly, the airway
downstream after the collapsed zone was permeable
and free of secretion.

The surgical procedure lasted 30 min. Haemoglo-
in oxygen saturation was maintained above 90% by
increasing FiO₂ and performing several recruitment
manoeuvres. After reversal of the neuromuscular block
and sevoflurane washout, the patient recovered sponta-
neous ventilation. SpO₂ quickly rose above 96%, and
lung auscultation returned to normal with symmetrical
vesicular murmur. At this point, the endotracheal tube
was still in place. It was removed once standard criteria
for extubation were met.

On re-reading the patient’s file after the procedure,
we noticed that a similar episode of desaturation had
already occurred during a previous anaesthesia but had
not been further investigated. Of note, no evidence of
extrinsic compression by an intrathoracic tumor, cyst,
aortic aneurysm, vascular anomaly or goitre was found
on recent chest X-rays and CT scans.

Fig. 1 Collapse of the left main bronchus
**Discussion and conclusions**

Physiologic narrowing of the central airway occurs during expiration. Weakness in the structure of the trachea and/or the bronchi may cause this narrowing to become excessive. A clear cut-off between physiological and pathological narrowing has yet to be determined [2]. When extraluminal pressure exceeds intraluminal pressure, as during expiration or coughing, the affected areas may then collapse. These conditions are referred to as ECAC [1, 3, 4]. The causes of ECAC in adults are listed in Table 1.

Non-invasive positive pressure ventilation is the cornerstone of treatment for ECAC. Positive pressure acts as a pneumatic stent, keeping airways open and improving respiratory flow. As a last resort, permanent mechanical stenting or surgical stabilization may be considered [1, 5].

If a patient with ECAC must undergo anaesthesia, spontaneous breathing should be maintained and CPAP applied. If not possible, PEEP should be applied during positive pressure ventilation. Airway collapse should be suspected in the following circumstances: hypoxia, increased insufflation pressures in volume control ventilation, and decreased tidal volume in pressure control ventilation. Recruitment manoeuvres should be attempted, and PEEP should be increased. Positional change should also be considered (reverse Trendelenburg or sitting position). If the obstruction persists, temporary mechanical stenting can be achieved by pushing the tracheal tube past the collapsed zone under fibroptic bronchoscopy control. Jet ventilation can also be attempted. As a last resort, mechanical circulatory support should be considered [6].

A case report previously described an adult patient with cerebral palsy presenting with acute desaturation during the induction of anaesthesia [7]. Haemoglobin oxygen saturation rapidly improved with positive pressure ventilation. However, in spontaneous ventilation (negative airway pressure), desaturation occurred again. In this case, the cause was extrinsic compression of the trachea and left mainsteam bronchus by the aorta.

Atkins et al. reported an 85-year-old patient with undiagnosed ECAC who was scheduled for Botox injections as a treatment for achalasia [8]. The patient began coughing during the procedure, and air flow stopped despite respiratory efforts, as assessed by paradoxical chest wall movements. Air flow was immediately restored by positive pressure ventilation. The main hypothesis was therefore that a collapse of the trachea occurred due to coughing.

In both cases described above, positive pressure ventilation improved gas flow and oxygen delivery. The particularity of the case reported here is the fact that the patient’s left main bronchus was permeable in spontaneous breathing but collapsed during general anaesthesia, including neuromuscular blockade. SpO2 increased slightly with recruitment manoeuvres (PEEP> 30 cm H2O for 40s), suggesting that the collapse was at least partially lifted. However, even during these recruitment manoeuvres, auscultation showed a lack of effective ventilation of the left lung, indicating that even with high pressures, normal flow could not be restored. Our hypothesis is that mechanical bending of the left main bronchus occurred, leading to its complete obstruction. In spontaneous breathing, the patient would maintain a particular body position (slightly curved on its left side) and a thoracic muscle tone, keeping the left main bronchus permeable. Under anaesthesia, including neuromuscular blockade, this natural conformation was likely lost, and the bronchus bent until it collapsed. One of the difficulties in diagnosing ECAC resides precisely in the fact that the degree of collapse can vary depending on the respiratory pattern and the patient’s position [1]. If the patient was to present for longer procedures, a double lumen tube could be used, with the left endobronchial tube bypassing the lesion. The placement of a temporary or even permanent stent could also be considered.

The fact that we were not able to verify our hypothesis is a limitation to this case report. One way to do this would be to perform general anaesthesia while keeping the patient in spontaneous breathing. Fibroptic

| Table 1 Causes of ECAC [1, 3] |
|--------------------------------|
| **Congenital**                  |
| • Tracheo-oesophageal fistula    |
| • Oesophageal atresia           |
| • Chromosomal defects           |
| • Tetralogy of Fallot           |
| • Genetic disorders             |
| **Acquired**                    |
| • Compression by great vessels (Brachiocephalic, aorta, pulmonary artery) |
| • Bronchopulmonary dysplasia    |
| • Pulmonary/mediastinal cysts    |
| • Intrathoracic tumors          |
| • Large/ectopic thymus          |
| • Goitres                       |
| • Heart/lung transplant         |
| • Liver failure/transplant       |
| • Prolonged intubation          |
| • Tracheostomy                  |
| • Emphysema                     |
| • Chronic bronchitis/obstructive pulmonary disease |
| • Relapsing polychondritis      |
| • Mucocutaneous leishmanias     |
bronchoscopy could then be performed to assess airway permeability while anaesthesia is deepened, and neuromuscular blockade induced.

In conclusion, applying positive pressure to the airways is the cornerstone of ECAC treatment. Interestingly, this approach has been unsuccessful in the case described herein. When possible, spontaneous ventilation should be maintained in patients with known or suspected ECAC.

Abbreviations
COPD: Chronic obstructive pulmonary disease; CPAP: Continuous positive airway pressure; ECAC: Expiratory central airway collapse; PEEP: Positive end-expiratory pressure.

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GG, L‑MN and PD treated the patient. GG and L‑MN wrote the manuscript. PD and SM revised the manuscript. All authors read and approved the final manuscript.

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Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
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Competing interests
The authors declare that they have no competing interests.

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