Successful Treatment of Cerebral Arterial Gas Embolism Following Uneventful TBNA

Monica Rocco1, Antonio D'Andrilli2, Alessandro Bozzao1, Luigi Maggi1*, Erino Angelo Rendina2 and Roberto Alberto De Blasi1

1Department of Surgical and Medical Science and Translational Medicine, Anesthesia and Intensive Care, Sapienza University of Rome, Italy
2Department of Surgical and Medical Science and Translational Medicine, Thoracic Surgery, Sapienza University of Rome, Italy

Abstract

Fibrobronchoscopy is commonly considered a safe procedure with a low major complication rate not including cerebral arterial gas embolism (CAGE) a severe life threatening iatrogenic complication. Several cases of transbronchial needle aspiration (TBNA) has been related with CAGE when patient happens to have the high airway pressure that exceeds the pressure of the pulmonary veins allowing the air to enter the systemic circulation through the left heart. HBOT is the only effective treatment available for CAGE that provides 100% oxygen at high pressure, which accelerates nitrogen reabsorption and improves oxygenation of ischemic tissue.

We reported a case of successful treatment with Hyperbaric Oxygen therapy of CAGE induced by an uneventful transbronchial biopsy during fibrobronchoscopy.

Keywords: Bronchoscopy; Trans bronchial needle aspiration; Hyperbaric Oxygen Therapy; Cerebral Arterial gas embolism

Background

Fibrobronchoscopy is commonly considered a safe procedure with a low major complication rate of 0.74% [1] that does not include CAGE [2], a severe life threatening iatrogenic complication occasionally reported during invasive procedures, including fibrobronchoscopy [3-5].

HBOT is the only effective treatment [6] available for Cerebral arterial gas embolism (CAGE) that provides 100% oxygen at high pressure, which accelerates nitrogen reabsorption and improves oxygenation of ischemic tissue.

We reported a case of successful treatment with Hyperbaric oxygen therapy (HBOT) of CAGE induced by transbronchial biopsy during fibrobronchoscopy.

Case Report

A 75-year-old man with history of hemoptysis underwent computed tomography (CT) scanning and was found to have two lung opacities with irregular margins involving the hilar portion of the right upper and middle lobes. Associated hilar and subcarinal enlarged lymph nodes were also found.

An outpatient flexible bronchoscopy showed a normal right bronchial tree, except for a mild restriction of the anterior segment branch of the upper lobar bronchus due to external compression. A Trans bronchial needle aspiration (TBNA) was performed at the level of the subcarinal and hilar nodes at the site of the mild bronchial restriction. Endo-bronchial ultrasound (EBUS) was not used. Procedures were easy and rapid to perform, and no significant endobronchial bleeding was observed. At the end of the procedures, the patient was unconscious with a Glasgow Coma Score of 4, but all brainstem reflexes were maintained with no hemodynamic or respiratory alterations. The patient was rapidly intubated and admitted to the intensive care unit where an episode of epilepsy occurred. Brain CT scan showed frontal right-sided cerebral air embolism (Figure 1a). High doses of oxygen and a bolus of 5 mg of midazolam followed by continuous infusion of 0.05 mg/Kg/h of midazolam were prescribed and the patient was transferred to the hyperbaric center.

The patient’s condition deteriorated and he was rapidly intubated and transferred to the intensive care unit. On the following day, the patient was virtually unconscious, with a Glascow Coma Score of 4, but all brainstem reflexes were observed. At the end of the procedures, the patient was unconscious and remained in this condition for 360 minutes. The day after HBOT, the patient recovered neurological signs (Glasgow Coma Score 15) and a brain CT scan showed a complete resolution of the cerebral air embolism (Figure 1b). On day 4, the patient was discharged from the intensive care unit. Cytological finding was: chronic inflammatory tissue with some “Actinomices” like colonies. Final diagnosis was a slowly resolving pneumonia.

Discussion

We reported a case of a severe CAGE following an uneventful fibrobronchoscopy and biopsies. Despite embolism not being included in fibrobronchoscopy guidelines as a major complication, CAGE associated with this procedure has been previously documented [5]. At the beginning of the twentieth century, Van Allen et al. [8] described two mechanisms for arterial gas embolism: arterialization of venous bubbles via a patent foramen ovale, and infusion into the pulmonary vein. There is an abundance of literature describing venous iatrogenic embolism resulting from many different procedures [9] that can produce cerebral symptoms in the presence of cardiac right to left shunts, or pulmonary arteriovenous malformation.

Other invasive procedures cause iatrogenic CAGE by introducing air directly into the left circulation [10]. Among these, transbronchial needle aspiration could cause a fistula between pulmonary veins and the airway [10] when the needle traverses both a pulmonary vein and an adjacent airway. If the airway pressure is elevated, as during cough in a semi-recumbent position, it exceeds the pulmonary venous pressure allowing the air to enter the pulmonary vein, reaching the systemic source are credited.

*Corresponding author: Luigi Maggi, Department of Surgical and Medical Science and Translational Medicine, Anesthesia and Intensive Care, Sapienza University of Rome, Italy, Tel: 00393895820203; Fax: 003933775400; E-mail: maggi.medicina@libero.it

Received October 27, 2015; Accepted December 14, 2015; Published December 18, 2015


Copyright: © 2015 Rocco M, et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.
circulation through the left heart.

There seems to be no relationship between the gas volume and neurological symptoms. Usually, there are small bubbles that reach 30-60 μm of diameter in the cerebral arteries, leading to ischemia and endothelial injury [11]. This dynamic process determines a wide variability in the clinical presentation and difficult diagnosis [12] with a possible delay in the treatment. In the case of suspected CAGE, it is currently recommended to administer pure oxygen and, thereafter, to refer the patient to a hyperbaric facility as soon as possible [6].

The high oxygen tension promotes the reabsorption of nitrogen from the bubbles and the elevated ambient pressure reduces their size in accordance with Boyle's law. At a pressure of 282 kPa, the spherical gas bubble diameter is reduced to 82% with a resulting 45% decrease in volume with resolution of embolic phenomena.

Weenick et al. [13] failed to demonstrate a positive effect of HBOT administered 2 or 4 hours after CAGE in pigs, but their experimental strategy caused very severe CAGE irresponsive to any HBOT effects and they avoided administering oxygen prior to HBOT. In a prospective study however, Bessereau et al. [14] reported that a longer time interval between CAGE and the first HBOT did not affect mortality but aggravated neurological sequelae at 1-year follow-up.

In our case, we reported an early hyperbaric treatment of a severe CAGE induced by fibrobronchoscopy. Full neurological recovery confirms the opinion of Souday [11] that CAGE is a race against time. Surely guidelines should provide awareness of potential CAGE complications after fibrobronchoscopy for a rapid diagnosis.

References