Anterior Spinal Artery Syndrome as Complication of Bronchial Artery Embolization

Ka Hong Chan MD, Chris White MD, FRCPC, and Jason K Wong MD, FRCPC

ABSTRACT
Bronchial artery embolization (BAE) has risen as one of the cornerstones of massive hemoptysis management. Though rare, spinal cord infarction is a potential complication. Here, we present a case of a 65-year-old gentleman who presented with acute weakness and was diagnosed with spinal cord infarction following BAE. This case will also review the pathophysiology of this adverse complication.

RESUME
L'embolisation de l'artère bronchique (BAE) est devenue l'un des piliers de la prise en charge massive de l'hémoptysie. Bien que rare, l'infarctus médullaire est une complication potentielle. Nous présentons ici le cas d'un homme âgé de 65 ans qui s'est présenté avec une faiblesse aiguë et un diagnostic d'infarctus médullaire à la suite d'une BAE. Ce cas examinera également la pathophysiologie de cette complication indésirable.
sided BA conducted subsequently continued to demonstrate abnormal neovascularity. Further embolization was conducted by inserting a microcatheter past some tiny branch origins, which was suspected to provide supply to the spinal cord (Figure 1).

The next day, the patient complained of left sided weakness and numbness on the belly. Strength examination demonstrated 4/5 weakness on the left lower as well as decreased pain and temperature sensation up to the T5-T6 area on the right. The rest of his physical examination, including reflexes and tone, were normal. Magnetic resonance imaging of his thoracic spine demonstrated a subtle linear T2/STIR hyperintensity within the anterior spinal cord at the T6-T8 level consistent with cord ischemia (Figure 2). Fortunately, his functional deficits were mild, and he made a good recovery with rehabilitative therapy.
Discussion

BAE has now become the mainline treatment for massive hemoptysis as many of these patients are poor surgical candidates due to their underlying chronic pulmonary disease (e.g., interstitial lung disease, cystic fibrosis, chronic infections such as tuberculosis). Identification of the BA is the cornerstone of this treatment. However, there is significant anatomical variability in the branching pattern and multiple possible anomalous origins of the BA. The diffuse involvement of these patients’ underlying pulmonary disease make the exact site of hemorrhage is difficult to confidently isolate. Therefore, because of the life-threatening nature of massive hemoptysis, multiple BA are frequently treated to reduce bleeding risk.

Radicular and medullary arteries, which reinforce blood flow to the anterior spinal artery, can arise from the BA as well as the intercostobronchial trunk and intercostal artery. Hence, a side effect when embolizing the BA can be spinal cord infarction. The latter is especially important, and hence, should be identified during angiography by looking for the characteristic “hairpin” configuration. In reality, spinal arteries are difficult to observe and increased risk of ischemia can occur with repeated embolization as with our case. Moreover, care must also be taken to ensure that the artery of Adamkiewicz is not found in the upper thorax prior to embolization. Although it is mostly found T8 or lower, it can be found as high as T5.

Great care and multiple angiographies were conducted in our case to visualize and avoid the spinal arteries. Despite this, our patient still suffered from anterior spinal artery syndrome post BAE. In reality, spinal arteries are actually rarely observed and increased risk of ischemia can occur with repeated embolization as with our case.

In conclusion, although BAE has now arisen as a relatively safe procedure with good long-term outcomes, spinal cord ischemia can be a rare but serious complication in which patients should be counselled on prior to the procedure.

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Ka Hong Chan: design of case report, drafting and revising the manuscript. Jason K. Wong and Chris White design of case report and revising manuscript.

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References