A therapeutic conundrum: recurrent cystic-fibrosis-related haemoptysis complicated by acute pulmonary embolism

William G Flight,1,2 Rowland J Bright-Thomas,1,2 Stephen Butterfield,3 Andrew M Jones,1,2 A Kevin Webb1,2

CASE REPORT
Dr Flight (WGF): A 72-year-old man with cystic fibrosis (CF) was admitted to hospital with recurrent haemoptysis. He was a compound heterozygote for the F508del and R117H-7T mutations of the cystic fibrosis transmembrane conductance regulator (CFTR) gene. CF had been diagnosed at the age of 65 prior to which he had been labelled as having asthma and non-CF bronchiectasis. Baseline forced expiratory volume in 1 s was 36% predicted and he was chronically infected with Pseudomonas aeruginosa and Burkholderia multivorans. His regular therapy included long-term oral corticosteroids for allergic bronchopulmonary aspergillosis and nebulised colistin. In addition to multiple complications of CF he also had a history of cataracts, age-related macular degeneration, hypertension, peptic ulcer disease, colonic polyps, benign prostatic hypertrophy and basal cell carcinomas of the skin.

On the day of admission he had expectorated approximately a third of a cup of bright red blood. Intravenous tranexamic acid was administered at a dose of 1 g four times daily alongside intravenous temocillin, tobramycin and co-trimoxazole. The haemoptysis gradually resolved and he was discharged 2 weeks later to continue oral tranexamic acid but this has not been resolved conclusively as yet.

WGF: Following discharge, the patient continued to experience streaks of blood in his sputum on a daily basis. He was re-admitted 2 weeks later with haemoptysis and symptoms of a viral upper respiratory tract infection. On the day of admission he had expectorated half a cupful of fresh red blood. On arrival to hospital he was comfortable at rest with normal vital signs. Oxygen saturations were 97% on air. An ECG revealed normal sinus rhythm and a chest radiograph was unchanged from previously.

Two hours after presentation, he suddenly developed breathlessness and chest tightness. Examination revealed him to be tachypnoeic, tachycardic and hypotensive. Oxygen saturations had dropped to 78% on air. A repeat ECG showed new right bundle branch block. An urgent CT-pulmonary angiogram demonstrated a large thrombus involving the left main, left upper and left lower lobe pulmonary arteries (see figure 1A).

Serum troponin I was elevated at 0.51 g/litre. This was consistent with a new left bundle branch block on the ECG and delayed pulmonary embolism, the patient was discharging to a chest clinic and was given intravenous tranexamic acid. The patient’s condition improved on the ward. The patient made a full recovery and was discharged 2 weeks later to continue oral tranexamic acid.
Chest clinic

Figure 1  (A) CT pulmonary angiogram demonstrating thrombus in the left main and upper lobe pulmonary arteries (white arrow). (B) Coronal CT reconstruction illustrating a dilated bronchial artery arising from the descending aorta (black arrow).

further bleeding must be avoided. When there is a risk of significant haemorrhage, anticoagulation is best achieved with intravenous unfractionated heparin and close monitoring of the activated partial thromboplastin time. In the event of life-threatening bleeding, the intravenous heparin can be quickly discontinued. If possible, the cause of the bleeding should be identified and reversed. In the event of bleeding preventing safe anticoagulation, the insertion of an inferior vena cava filter is a further therapeutic option.

WGF: Intravenous heparin was commenced and the patient was also treated with intravenous temocillin and tobramycin. High-flow oxygen was administered to achieve target saturations of 94–98%. Fluid resuscitation was given and tranexamic acid was discontinued. An inferior vena cava filter was considered but was not pursued as the CT examination showed no evidence of venous thrombosis in the legs or pelvis. Despite the above measures, the patient experienced ongoing substantial haemoptysis. Advice from the interventional radiology service was sought.

Dr Butterfield (Interventional Radiology): My first step was to review the original CT pulmonary angiogram study. CT reconstructions were performed which revealed a dilated bronchial artery arising from the descending aorta (see figure 1B). We elected to perform a bronchial artery angiogram with the aim of embolising any potential sources of bleeding from the bronchial circulation. The dilated bronchial artery identified on the CT scan was supra-selectively catheterised and embolised to stasis with polyvinyl alcohol.

The available evidence suggests that BAE is an effective treatment for massive haemoptysis in CF. There have been no randomised trials of BAE in CF but large case series suggest that acute bleeding is controlled in up to 97% of cases with a long-term recurrence rate of 27–46%.5 Chest pain is the most common side effect following BAE, although potential complications include an acute neurological deficit due to embolisation of a spinal artery or passage of embolic material through collateral vessels to the cerebral circulation. In the present case the patient experienced mild chest pain for 24 h after the procedure but did not suffer any more serious adverse events.

WGF: Following BAE the patient’s haemoptysis settled completely. Intravenous heparin was converted to low molecular weight heparin and oral warfarin was commenced 48 h after undergoing BAE. He was discharged at day 16, by which time his symptoms had returned to baseline and he had remained free from haemoptysis.

AKW: The development of PE in a patient with ongoing major bleeding presents a highly challenging clinical situation that may be encountered in a wide variety of medical and surgical specialties. The principles of management in this situation are applicable to a number of conditions in respiratory medicine, such as the patient with lung cancer, aspergilloma or myriad other causes of haemoptysis.

The scenario described is highly unusual in CF however, and to my knowledge has not been reported before. Idiopathic PE alone is considered exceptionally rare in patients with CF. Previous reports of PE in CF have all been associated with venous access device related thrombosis. In this case it is certainly possible that the use of the tranexamic acid contributed to the development of venous thrombosis. The risks associated with such pro-thrombotic drugs remain uncertain and need to be addressed in future studies.

As the CF population ages it is reasonable to expect that the incidence of venous thrombosis in this population will steadily rise. Since haemoptysis is such a common feature of CF, it stands to reason that we shall also see concurrent haemoptysis and thromboembolic disease with increasing frequency.

The key learning points to be taken from this case are:
1. The CF population is getting older and diseases associated with ageing are increasingly relevant to the CF physician.
2. Bronchial artery embolisation represents the therapy of choice for massive haemoptysis in CF.
3. CF physicians should be alert to the possibility of PE in their patients, a diagnosis which is easily overlooked in the context of severe CF lung disease.
4. The conundrum of acute PE in the context of major bleeding requires a delicate balance of anticoagulation and correction of the underlying haemorrhage.

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