CASE BASED DISCUSSION

A lesson in plasticity: a 74-year-old man with plastic bronchitis

Anand Shah,1,2 Jackie Donovan,3 Phil Marino,4 Pallav L Shah,1,2 Anand Devaraj,2,5 Kshama Wechalekar,2,6 Toby M Maher,2,7 Michael R Loebinger,1,2 Robert Wilson1,2

CASE BASELINE DISCUSSION

AS: A 74-year-old male was transferred to the Royal Brompton hospital from a local hospital with respiratory failure and expectoration of bronchial casts (see figure 1A). The patient recalls recurrent chest infections with occasional haemoptysis in his teens with no prior history of asthma, TB or pertussis. Despite investigation and antibiotic treatment, his symptoms of intermittent, chronic recurrent bronchial cast expectoration continued. A fibroptic bronchoscopy on the patient, aged 41, revealed erythematous bronchial mucosa in the right middle lobe, with a ventilation/perfusion scan revealing concomitant decreased ventilation. Bronchial washings were unremarkable, and given concern of a pathology originating from the right middle lobe, a right middle lobectomy was performed. Histology showed non-specific inflammatory fibrosis around the lobar bronchi.

His symptoms, however, did not resolve. Microbiological examination of his casts revealed no growth, and histological examination showed fibrin and mucus with some lymphocytes. Further trials of corticosteroids, nebulised N-acetylcysteine, hypertonic saline, prophylactic azithromycin, systemic antifungals and oral cyclophosphamide were unsuccessful. Extensive immunological and allergic tests were normal.

His symptoms remained stable till age 73 when cast expectoration and infections grew more frequent. Recently, his condition had deteriorated, requiring admission locally with respiratory failure. He was transferred to our centre for further evaluation.

On arrival, the patient was in respiratory distress. A FiO2 of 60%, oxygen saturations were 90% with arterial blood gas analysis showing a pO2 of 6.72 kPa, pCO2 of 6.31 kPa, pH of 7.38 and bicarbonate of 30.5 mmol/L. A chest radiograph showed significant right-sided consolidation. Blood tests revealed a normal white cell count with a C reactive protein of 10 mg/L. He was empirically started on intravenous tazocin, caspofungin, hydrocortisone and octreotide causes mild vasoconstriction of splanchinic vessels, including hepatic venous outflow. This leads to reduction in gastric, pancreatic and intestinal secretions as well as intestinal absorption collectively reducing the flow of chyle. The patient, however, had significant bronchial casts, and a means of either removal or liquefaction was required. A flexible bronchoscopy approach was considered.

PS: Removal of bronchial casts by bronchoscopy is difficult. The casts are often too firm for removal
by suction, yet too friable for removal by forceps. Novel technology such as cryotherapy may be useful, and hence bronchoscopy was attempted. No bronchial casts were visible, however, with oedematous collapsed right lower and upper lobe bronchi. No therapeutic procedures were performed. Microbiology was unrevealing, and bronchial biopsies showed non-specific inflammation.

AS, ML and RW: We hence focused on cast liquefaction. The use of both nebulised dornase-α (DNase) and tissue plasminogen activator (t-PA) has been reported for the management of chylous bronchial casts. Due to coexistent dual antiplatelet therapy following drug-eluting coronary arterial stent implantation, we initially examined in vitro efficacy.

JD: A dried bronchial cast expectorated by the patient was incubated for 2 hours in 2 mL of 0.1 mg/mL t-PA, 0.1 mg/mL DNase or saline. The cast was removed, air-dried and reweighed, with supernatant analysis for protein and fibrin breakdown products. Incubation with t-PA resulted in structure loss, 62% cast mass reduction, 0.47 g/L protein and >4×10⁶ ng/mL soluble cross-linked fibrin (XDP) release. DNase incubation resulted in no structural change, 10% mass reduction, 0.07 g/L protein and 40 000 ng/mL XDP release. Saline incubation resulted in no structural change, 2% mass reduction, 0.0 g/L protein and 1200 ng/mL XDP release.

AS, ML and RW: Biochemical analysis showed partial and high liquefaction efficacy of DNase and t-PA, respectively. Within a high-dependency environment, nebulised DNase 2.5 mg, two times per day, and t-PA 5 mg, three times per day, were started. No haemorrhagic complications were seen with radiological improvement within 24 hours. However, liquefaction led to difficulty in clearance, intermittent hypoxia and cast reformulation with further bronchial obstruction. We therefore focused on understanding the pathophysiology of chylous bronchial cast formation.

PM: Post Fontan procedure due to congenital heart disease, it is thought that high central and pulmonary venous pressures may play an aetiological role in cast formation. Hence, an echocardiogram was performed. This showed normal left ventricular systolic and diastolic function with mild hypokinesia of the posterior inferior wall. The right heart was mildly dilated with normal longitudinal function. There was no significant valvular disease, normal-sized right atrium and an estimated pulmonary arterial systolic pressure of 51 mm Hg. Given evidence of right heart strain on echocardiogram and moderately elevated pulmonary pressures, the patient was started on pulmonary vasodilator therapy with sildenafil and uptitrated to 50 mg three times per day.

AS, ML and RW: There have been recent reports regarding the use of the mammalian target of rapamycin (mTOR) inhibitor sirolimus (rapamycin) for management of recurrent chylothoraces in patients with lymphangioleiomyomatosis.

TM: Lymphangioleiomyomatosis is a progressive cystic lung disease in women associated with inappropriate activation of the mTOR signalling pathway causing dysregulation of cellular growth and lymphangiogenesis. This can result in recurrent chylothoraces and, on rare occasions, chyloptysis. Recent studies have highlighted the successful use of sirolimus in the management of chylothoraces and stabilisation of lung function. The mechanism of action is unclear, but potentially relates to the regression of abnormal smooth muscle cellular proliferation.
causing obstruction and leak within the lymphatic system. In the current case, the reported symptoms since childhood may point to a diagnosis of generalised lymphatic anomaly.

Generalised lymphatic anomaly is a group of disorders that can include conditions with congenital, benign, focal areas of lymphatic proliferation occurring in any part of the body containing lymphatics. This pathological process can result in restrictive and/or obstructive physiology, chylous effusions and respiratory failure and is predominantly detected in children and young adults.

AS, ML and RW: Given the likelihood of lymphatic obstruction alongside clinical deterioration, we started treatment with sirolimus aiming for a therapeutic range of 5–10 ng/mL. Gradually, the patient improved and was discharged from hospital. Subsequent lung function and CT imaging revealed almost complete resolution of his right-sided lung changes and markedly improved lung function. To confirm lymphatic obstruction, a lymphoscintogram was performed (figure 1E). This showed abnormal activity predominantly in the right lung, confirming chylous leak. Recently described novel magnetic resonance lymphangiography techniques potentially enable improved imaging and therapeutic percutaneous embolisation of thoracic lymphatic leak. This expertise however is limited and was unavailable locally at the time of the patient’s presentation.

Three months after discharge, the patient no longer expectorated bronchial casts and continued on sirolimus, one time per day DNase and a low-fat medium chain triglyceride (MCT) diet. Two years later, only rare small episodes of chylous bronchial cast expectoration have occurred with a dramatic improvement in quality of life and respiratory function. No significant side effects from sirolimus therapy have been noted.

AS: Sirolimus is a promising therapeutic option in the management of patients with lymphatic obstruction or chylous leak. It is a macrolide structurally related to the calcineurin inhibitor FK506 and has potent immunosuppressive properties. Sirolimus binds with and inhibits mTOR, a serine/threonine kinase regulated by phosphoinositide-3-kinase. This inhibits the mTOR-mediated signal transduction pathways, resulting in the arrest of cell cycle in G1 phase in various cell types. Inhibition of interleukin 2-induced proliferation of T cells also makes it a valuable immunosuppressant to prevent organ rejection in solid-organ transplantation. Recent genetic studies in patients with isolated and syndromic lymphatic malformations, however, have shown somatic mutations in the catalytic subunit of phosphoinositide-3-kinase (PIK3CA) causing overproliferation of lymphatic endothelial cells with improvement on sirolimus therapy with antilymphangiogenic effects. This finding may explain the underlying mechanism of action, and importantly, provide a future target for gene therapy in congenital lymphatic obstruction.

Contributors All authors were involved in the clinical care of the patient in this case. JD provided biochemical analysis. AD and KW provided radiological input. AS drafted the manuscript, and all authors assisted in the editorial process.

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REFERENCES