Recurrent pulmonary embolism related with Paget–Schroetter syndrome: a case report

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Received 28 December 2018; first decision 21 March 2019; accepted 21 June 2019; online publish-ahead-of-print 18 July 2019

Background
Paget–Schroetter syndrome (PSS) is an unusual cause of venous thromboembolism, which is frequently misdiagnosed and undiagnosed in clinical settings. Although axillary-subclavian vein thrombosis is related with PSS typically presents in healthy young athletes, it is possible for this phenomenon to occur in various age settings.

Case summary
We present a case of recurrent pulmonary embolism caused by a thrombus in dilated axillary vein related with PSS. A 74-year-old man was referred to our cardiology department for chest discomfort and hypoxaemia. The contrast computed tomography (CT) revealed that he suffered from bilateral pulmonary embolism. However, we could not find the source of embolism despite other examinations such as ultrasonography of the inferior limb deep vein. Three months later, the patient complained of dyspnoea for a second time, and a contrast CT scan was subsequently performed revealing a new pulmonary embolism. Surgical resection of the giant thrombus was performed, resulting in a good clinical course without recurrence.

Discussion
We experienced a case of recurring pulmonary embolism in a patient with undiagnosed PSS, which was related to the active and vigorous movement of the right arm during his working. Although there are various treatments for PSS including anticoagulation, first rib resection, and lifestyle modification, we need to consider what is the best treatment individually.

Keywords
Case report • Pulmonary embolism • Paget–Schroetter syndrome • Thrombosis

Introduction
Upper extremity deep vein thrombosis (UEDVT) is known as a rare diagnosis and currently accounts for ~10% of all deep vein thrombosis (DVT) cases. Paget–Schroetter Syndrome (PSS), also known as effort thrombosis, is an unusual cause of UEDVT and has remained frequently undiagnosed or misdiagnosed. We report a case of PSS, resulting in recurrent pulmonary embolism.
Case presentation

A 74-year-old man working as a car washer visited our cardiology department for chest discomfort and hypoxaemia. He had no previous medical history and was not taking any medications. On arrival, his temperature was 36.8°C, his blood pressure was 94/76 mmHg, his pulse rate was 117 beats/minute (regular rhythm), his respiratory rate was 24 breaths/minute, and his oxygen saturation was 89% on room air. Despite the presence of hypoxaemia and tachypnoea, we could not detect any other abnormal findings from his physical examination suggesting any disease related to hypoxaemia. Electrocardiogram on admission suggested right ventricular strain pattern with complete right bundle branch block and S1Q3T3 pattern. Echocardiography on admission revealed that remarkably enlarged right ventricular suppressed left ventricular contraction (Figure 1A). Hence a contrast CT scan was performed, revealing bilateral pulmonary embolisms to be the underlying cause of the symptom (Figure 1B). Despite other examinations, such as ultrasonography of the inferior limb deep vein, the source of the thrombosis could not be found. Laboratory findings revealed congenital coagulation disorders, including protein C/S deficiency, anti-thrombin III deficiency, and anti-phospholipid antibody syndrome to be negative.

After admission, the patient was initially administered systemic thrombolytic therapy with monteplase (1,200,000 unit) and heparin. Contrast CT scan performed 7 days after admission revealed complete disappearance of the thromboembolisms and symptoms were mitigated as well.

Three months later, the patient was referred to our hospital because of recurring chest discomfort. Another contrast CT scan was performed, revealing a new pulmonary embolism despite continuous anticoagulation therapy with warfarin. PT-INR during treatment period was in the standard treatment range [PT-INR (1.6–2.6)] and PT-INR on second admission was 2.1. A giant thrombus (43 × 50 mm) located in the right axillary vein, which we could not find during previous hospitalization, was pointed out (Figure 1C). Vein angiography was also performed, revealing a giant thrombus with no collateral vessels. Since ultrasound echography showed sufficient blood flow around the thrombus, swelling, and pain of his right arm were not observed before the operation.

We were finally able to identify that the source of his recurrent pulmonary embolism was a giant thrombus located in the right dilated axillary vein. Eventually, the huge thrombus was removed surgically (Figure 1D), and the symptoms were subsequently alleviated. Anticoagulation with warfarin was continuously prescribed and chest discomfort completely resolved. The patient retired his job due to this experience and his old age, and reported no post-operative complication during 2 years after the operation.

Discussion

Upper extremity deep vein thrombosis is classified into primary UEDVT and secondary UEDVT; most cases of primary UEDVT are iatrogenic (inserting central venous catheter, pacemaker insertion, and parenteral nutrition) whereas secondary causes are often due to hypercoagulable state and surgeries involving the upper limbs.

Paget–Schroetter syndrome, also referred to as ‘effort thrombosis’, usually occurs in young athletes such as tennis players, judo players, and those who use their dominant hand and arm. Paget–Schroetter syndrome remains a relatively rare condition; it occurs at a rate of 2 per 100,000 people per year.² It is believed that active and sustained movements during sports lead to microtrauma of the endothelium and activation of the coagulation cascade. According to some reports, the anatomical abnormalities involving the thoracic outlet play an important role in the pathogenesis of making microtraumas of endothelium.³,⁴

The anatomical abnormalities involving the thoracic outlet (cervical rib, congenital bands, hypertrophy of scalenus tendons, and abnormal insertion of the costoclavicular ligament) are important regarding the pathogenesis of effort thrombosis.³,⁵ The narrow costoclavicular space leads to continuous compression of the vein and restricts blood flow, especially when the arm is hyperabducted or externally rotated. Therefore, it is considered that PSS is a sequel of thoracic outlet syndrome (TOS).
Chronic compression of the vein leads to an inflammatory response in the soft tissues surrounding the vein and endothelial micro-trauma. The repetitive endothelial trauma results in intimal hyperplasia, inflammation, and eventually hypercoagulable state. In the present case, it can be considered that vigorous right arm abduction during car washing led to chronic compression of vein.

As there are only few reports mentioning thrombo-embolic pulmonary hypertension induced by UEDVT, the optimal therapeutic approach has not been well defined. However, since residual deep vein thrombosis will become the cause of recurrent pulmonary embolism, chronic thrombo-embolic pulmonary embolism, and disabling symptoms, aggressive treatment for both DVT and hypoxaemia are necessary. Some authors suggested that the surgical decompression of the thoracic outlet is a necessary part of the treatment since PSS is considered as a sequela of TOS. The removal of the first rib through a transaxillary approach is commonly performed. However, in the present case, recurrence of PSS was not observed during follow-up period without a surgical decompression of the thoracic outlet.

**Conclusion**

We reported the case of undiagnosed and repeated pulmonary embolism related with PSS. Not only lower extremity but also upper extremity vein should be considered as the source of embolism of pulmonary embolism.
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