Primary Intimal Pulmonary Vein Sarcoma with Expansion to the Left Atrium

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Received: July 11, 2017; Published: July 18, 2017

Abstract

We have recently treated a patient with pulmonary vein sarcoma. It is a 65 year old women who was admitted to the hospital because of the chest pain with propagations down her right arm, followed by suffocation and coughing out content with traces of blood and febrility >38°C. The primarily suspected pulmonary embolism was ruled out after the diagnostic heart CT scan and transthoracal and transoesophageal echocardiogram which verified the existing of a tumor mass in the left atrium. The patient underwent an urgent surgery and the tumor mass was removed surgically from the upper right pulmonary vein. The path histological diagnoses revealed pulmonary vein intimal sarcoma. Pulmonary vein intimal sarcoma is one of the rarest sarcoma subtypes. To our knowledge, there is only one published case of such pathology in the literature.

Introduction

A 65 year old female patient was admitted in the Emergency Room of Montenegrin Clinical Hospital because of the chest pain with propagations down her right arm, followed by suffocation and coughing out content with traces of blood and febrility >38°C. She mentioned the poor physical exercise tolerance with occasional swelling of lower extremities. She was experiencing the discomforts for the previous 10 days [1-3]. The patient had been a smoker for a long time. She was aware of cardiac arrhythmia which is why she had been prescribed an oral anticoagulant therapy. She had been aware of her condition of thrombocytopenia too. Twenty years ago she underwent hysterectomy with bilateral adnexectomy.

During the physical examination the patient was eupnotic while resting, acyanotic, anicteric and afebrile. Her vital signs were: cardiac frequency (FR) 85/min, blood pressure (TA) 100/60 mmHg, respiratory frequency (FR) around 16-18/min, saturation (SO2) around 96%. Pulmonary auscultation revealed a weakened basal respiratory wheeze with sporadic late inspiratory bilateral basal crackles [4-7]. The auscultation also showed a rhythmical cardiac activity with dear tones and without wheezes. The extremities were without edemas, varicosity or deformities. EKG showed the sinus rhythm of around 85/min, normogram, without signs of ischemia.

Laboratory Test Results

Se 30...72, CRP 10...160, D-dimer 0,73, PV 14,2, INR 1,2, fibrinogen 81, Le 5,74, Er 3,96, Hgb 122, HCT 0,35, Tr 162. Pulmonary X- ray showed left basal pleural pericardial effusion. Chest CT scan describes a 35 mm thrombus in the main part of Art. Pulmonalis which extends to the branch for the upper lobe with consequential parenchyma consolidation ventrally bigger than 50 mm, pericardially along with the pericardium almost completely occupying the front FC sinus and dorsally with the lower branch of interlobar incisura up to 30 mm and medially almost at the very top of the lungs with the size of up to 20 mm with the complete hypo perfusion of the upper lobe with evidently positive air bronchogram.

Heart CT scan describes in the left atrium a big tumor-like change with the dimensions of 32x24mm, 32 mm long along which thrombotic masses are evident. The tumor covers both lower left
Several tissue fragments with the size up to tissue. Path histological diagnosis intimal sarcoma (gradus 2). Mediastinal lymph node is of preserved structure, without tumor S100, CD 34 and HMB 45 negative. The path histologically analyzed Immuno histo chemical atypical spindle and oval cells. The nuclei are slightly to moderately in slime.

Patho Histological Results

Macroscopic: Several tissue fragments with the size up to 40mm and with softer to medium hard consistency, partly covered in slime.

Microscopic: Tumor tissue made of groups and solid clusters of atypical spindle and oval cells. The nuclei are slightly to moderately polymorphic, oval and hyper chromatic.

A. A moderate number of mitoses – focally between 5 and 10 mitoses on 10 fields of big enlargements.

B. The tumor contains bleeding zones and small necrosis focuses.

C. Hypo cellular slimy degenerated areas were present.

Immuno histo chemical

Tumor cells are Vimentin and Actin positive; CK, EMA, Desmin, S100, CD 34 and HMB 45 negative. The path histologically analyzed mediastinal lymph node is of preserved structure, without tumor tissue. Path histological diagnosis intimal sarcoma (gradus 2).

Discussion

Primary neoplasm of big blood vessels (aorta, pulmonary artery and pulmonary veins, v. cava sup and inf) are extremely rare. The pathogenesis of those tumors is still vague and signs and symptoms are unspecific which makes differential diagnosis difficult and postpones a final diagnosis. Around 400 cases of primary tumors have been mentioned in reference books so far, 30 of them being cases of primary pulmonary vein tumors. The majority of those tumors are leiomyosarcomas and there is one case of myxosarcoma and one case of myxoidfibrosarcoma. These pulmonary vein sarcomas are often located in the upper right and lower left pulmonary veins. To our knowledge this is the second published case of pulmonary vein non-myxoind intimal sarcoma (fibro sarcoma). The biggest group of patients with pulmonary vein primary sarcoma includes 17 cases of leiomyosarcoma. Blood vessel tumors appear on average when people are in their 40s and they are equally found in both sexes (leiomyosarcomas mildly predominate with females). The most common symptoms like cough, hemoptysis, dyspnea, chest pain and pleural effusions are unspecific. Pulmonary vein sarcomas, especially intimal sarcomas can be misdiagnosed as pulmonary thromboembolism, which was initially the case with our patient, because of the fact that it is rare and because of unclear clinical progress. The consequences can be inadequate treatments like a prolonged anticoagulant therapy or thrombolysis. In the biggest series of pulmonary vein leiomyosarcomas, around 50% of the cases were diagnosed with the help of CT and/or MRI angiography. It is important here to be mentioned that in majority of such cases the tumor masses are thought to be originating from left atrium. Bronchoscopy and transesophagheal echocardiography together with heart CT are important for proper diagnosing and urgent management of those cases.

The difficulties in diagnosing are only one of the aspects related to bad prognosis of primary pulmonary vein tumors, regardless of the subtype. Only two cases were recorded when patients lived three or more years after the treatment of leiomyosarcoma and only one patient lived more than nine months after the treatment of myxosarcoma. Both patients with leiomyosarcoma had a complete surgical resection with clean remaining edges. After the resections there was lots of case of relapsing which makes us conclude there is a need for adjuvant or non-adjuvant therapy. Surgery with adjuvant chemotherapy and radiotherapy can improve short term survival. However the prognosis is not good with the rate of five year survival between 0 and 6 percent.

The ideal management of pulmonary vein sarcomas is still being discussed. Extensive surgical resection is for now considered to be the most optimal solution. For patients with incomplete resection or local relapsing heart transplanting is taken into consideration as a valid solution. Although some authors do not think that chemotherapy and radiotherapy are efficient, some other authors hope that systemic chemo and radio therapy could increase the survival rate and achieving therapy and palliative benefits could avoid potential surgical complications.
References


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