DEATH DUE TO CARDIAC ANGIOSARCOMA: AUTOPSY CASE REPORT

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Background. Primary tumors of the heart are rarely detected at autopsy, especially angiosarcomas which are primary malignant one.
Objective. We presented autopsy case of cardiac angiosarcoma with morphologic findings.
Methods. We described adult man died in emergency service of the hospital.
Results. Reported case was 33 year-old-man who was died in emergency service of hospital where he was taken when he was ill after leaving home. According the prosecution documents, and the expressions of family, it was reported that he had a heart disease; his symptoms repeated 3 day ago before he died, he thought to attend the Cardiology Clinic due to his symptoms. At autopsy on macroscopic internal examination, mass with rough surface in the right atrium, hematoma at the posterior of the right atrium, blood in the pericardia, nodular lesions in hemorrhagic appearance in the sections of lung, liver and spleen were detected. In histopathologic examination; in the heart angiosarcoma as primary malign heart tumor and metastatic masses in the liver, spleen and lung were detected.
Conclusions. We aimed to discuss cardiac angiosarcoma case with autopsy and histopathologic findings in the aspect of medico legal literature.
KEYWORDS: Angiosarcomas; heart; metastatic tumors; autopsy.

Introduction
Primary tumors of heart are rare and they are revealed with prevalence between 0.0017 and 0.19 percent at autopsy [1]. While 75 percent of primary tumors of heart are benign, 25 percent are malign tumors [2]. Angiosarcomas, which are 35-40 percent of primary malign tumors, are the most common [3, 4]. Primary tumors are mostly developing in the right atrium and pericardium [5]. Diagnosis of patients is delayed until tumors become untreatable and many systemic metastases are present [4, 6]. The research is aimed to study the autopsy and histopathologic findings of the case in medical and legal aspect.

Case Report
The case involves a 33 year-old-man reported to pass away in the emergency department of hospital, where he was admitted, when he got ill after leaving home. In the analysis of the prosecution documents and according to the words of family members, it was reported that he had a heart disease; chest pain repeated 3 days before he died, he thought to attend the Cardiology Clinic due to the symptoms. In autopsy external examination, there were no pathologic findings, except for ecchymosis due to catheters on the dorsum of the right and left hands and left inguinal line. In internal examination revealed mass on the rough surface of the right atrium (Fig. 1), hematoma at the posterior of the right atrium, 500 cc free blood in pericardia, inhemorrhagic nodular lesions in the sections of lungs, liver and spleen (Fig. 2).

Histopathologically, in sections of heart, tumoral proliferation of atypical cells, which had a high mitotic index and spindle-oval nucleus and infiltrates between myocardial fibers, were present [4, 6]. The research is aimed to study the autopsy and histopathologic findings of the case in medical and legal aspect.

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Primary tumors of the heart are rare and are revealed with prevalence between 0.0017 and 0.19 percent at autopsy [1]. Approximately three-quarters of primary heart tumors are benign with atrial myxomas comprising three-quarters of them [2]. Angiosarcoma is the most common of malignant cardiac tumors [7]. Cardiac angiosarcomas can be found in right atrium, left atrium, pulmonary artery, but most commonly in the right atrium [8], as in our case.

Angiosarcoma is established to be as two-fold in men, typically between the third and fifth decade [4, 10] and our case is consistent with the literature. A study reported that a small series of heart sarcomas were subjected to a specific molecular study that proved the presence of K-ras mutations at codon 13 [11]. In human study mutation of the p53 tumor suppressor gene was present in cardiac angiosarcomas [12]. Cardiac sarcomas are rare but lethal disease. It is associated with a poor prognosis and median survival time from 1 to 81 months [6]. Angiosarcoma is associated with a high rate of hematogenous metastases, most of this tumors produce systemic metastases at the time of its detection, most commonly to the lungs, in addition to the liver, brain, adrenal glands, and bone [6, 5, 13]. In our case lung, liver metastases were present compliant with the literature. In addition, in our case there were splenic metastasis different from the literature.

Patients often have non-specific symptoms. So, it is difficult to diagnose cardiac angiosarcoma. Initial findings may include chest pain, dyspnea, fatigue, cough, heart murmur of unclear mechanisms, superior vena cava syndrome, constitutional symptoms, pericardial effusion, pericardial tamponade, arrhythmias, non-specific ST changes on ECG [6, 13, 14].

Echocardiography, magnetic resonance image (MRI) and computed tomography (CT) are used for diagnosis and systematic evaluation of cardiac angiosarcoma [15, 16, 17]. Echocardiography, transeosophageal or transthoracic, is the main tool for describing the localization and size of a mass [18]. There is no specific tissue density that is why the diagnosis of angiosarcoma is challenging. The histological examination of fluid taken by pericardiosentesis, revealed malignant cells. The definitive diagnosis of angiosarcoma is biopsy [15, 16]. Several studies established that infiltration of the myocardium by spindle cells, hyperchromatic nuclei, mitotic figures in various stages, giant cells, including tumor necrosis and hemorrhage in common areas, negative for cytokeratin, vimentin and FVIII were revealed as positive staining by histopathological examination [6, 15, 16]. Ge et al. reported that the tumor cells in cardiac angiosarcomas were strongly positive for CD31, CD34, FLI-1, and WT-1 but negative for AE1/3, D2-40, human herpesvirus 8, and epidermal growth factor receptor by immunohistochemical method [6].

The rareness of this disease limits gaining experience for adequate treatment. The therapeutic approach for primary cardiac angiosarcoma is surgery, chemotherapy and radiotherapy, alone or in combination. The initial treatment is surgery [6, 19].

Conclusions

Primary tumors of the heart are mostly originating in the right atrium and rarely observed at autopsy. Angiosarcomas are most commonly seen primary malign tumors. We discussed cardiac angiosarcoma case with autopsy and histopathologic findings in the aspect of medico legal literature.

Conflict of interest

The authors declare no conflict of interest.
РАПТОВА СМЕРТЬ ВНАСЛІДОК АНГІОСАРКОМИ СЕРЦЯ: РЕЗУЛЬТАТИ АУТОПСІЇ

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Вступ. Первинні пухлини серця вкрай рідко зустрічаються при аутоопсіях, особливо ангіосаркоми, які належать до злоякісних новоутворів.

Мета роботи привернути увагу та ознайомити широке коло читачів з результатами аутоопсії та морфологічними даними при ангіосаркомі серця.

Методи. Описано та проаналізовано результати аутоопсії дорослого чоловіка, який помер у відділенні невідкладної допомоги.

Клінічний випадок. 33-річний чоловік помер відділенні невідкладної допомоги, куди був доставлений з дому. Зі слів рідних у нього була хвороба серця, а перед смертю протягом трьох днів турбував біль у серці, з приводу якого він планував звернутися у лікарню. При аутоопсії виявлено новоутвір з шорсткою поверхнею у правому передсерді, гематому на задній стінці правого передсердя, 500 мл крові у перикарді, також було виявлено вузлові утвори при розтині легень, печінки та селезінки. За результатами гістологічного дослідження встановлено діагноз ангіосаркоми серця, як первинної пухлини, з метастазами у легені, печінку та селезінку.

Висновки. Описано рідкісний клінічний випадок ангіосаркоми серця, виявлений посмертно на аутоопсії.

КЛЮЧОВІ СЛОВА: ангіосаркома; серце; метастази; аутоопсія.

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