Rare Cause for Sudden Right Heart Failure

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Right heart failure occurs daily in clinical settings, but an underlying cardiac malignant tumor is very uncommon. We report a case of a 48-year-old man presenting only with palpitations and decompensated heart failure. Echocardiographic imaging revealed a large tumor of the right ventricle. Shortly after a putatively successful surgical approach, the patient was admitted again with heart failure symptoms. On reassessment, a complete relapse with multiple metastases could be seen. Generally, cardiac malignant tumors are diagnosed at a time-point when therapeutic options are very limited or even postmortem. Broad echocardiographic screening in patients with unspecific symptoms might be helpful to detect cardiac malignant tumors at early stages.

KEYWORDS: right heart failure, primary cardiac carcinoma, rhabdomyosarcoma, echocardiography

CASE

A 48-year-old, previously healthy man was admitted with acute decompensated right heart failure (RHF) causing massive lower extremity edema and dyspnea. The medical history was unremarkable except for paroxysmal cardiac palpitations for several months.

The cause for the acute right heart decompensation was identified in echocardiographic studies. Transthoracic and transesophageal echocardiography (TTE and TEE) revealed an obstructive structure embedded only within the right ventricle (RV), indicating a cardiac tumor (Fig. 1, Panel A; arrow). The complete staging did not identify other organ manifestations. Consequently, the tumor was removed surgically to eliminate the obstruction and to take specimens (Panel B). The subsequent pathologic assessment displayed a mainly undifferentiated spindle cell rhabdomyosarcoma. Consecutively, adjuvant chemotherapy was considered. The patient’s condition improved rapidly and he was discharged 6 days after surgery.

Three weeks after cardiothoracic tumor resection, the patient was admitted again with recurrent RHF symptoms, prior to initiation of chemotherapy. Reassessment unfortunately revealed a large tumor, similar to the previously known sarcoma, in the lateral RV (Panel C). Contrast-medium computed tomography (CT) as well as total body positron emission tomography (PET) with 18F-fluoro-2-deoxy-D-glucose (FDG) identified a complete relapse (Panels D and E)[1,2]. Due to physiological cerebral metabolic overlapping, PET only showed pulmonary FDG-avid lesions, while CT detected additional
multiple brain metastases. Considering this devastating cancerous progression and the patient’s wish, we discharged him to palliative care.

This striking case report impressively highlights that diagnosis of primary cardiac neoplasms is often made at the advanced stage because of the nonspecific nature of the symptoms that may suddenly exacerbate. Presentation of sudden illness with cardiac and constitutional symptoms or signs of systemic embolization may be susceptible for cardiac neoplasms[3,4]. Primary cardiac tumors are rare entities, but high percentages (~25%) of these are malignant tumors, mostly undifferentiated carcinomas, angiosarcomas, leiomyosarcomas, and rhabdomyosarcomas[5,6,7]. Larger prospective studies concerning management and treatment are missing[6]. The outcome in patients with malignant cardiac tumors is poor due to the limited therapeutic options and the late clinical manifestation, as demonstrated in the present case. The median survival is less than 1 year after initial diagnosis of a primary malignant cardiac tumor[5,6].

In contrast to primary cardiac tumors, metastatic involvement of the heart is over 20 times more common and has been reported in autopsy series in up to one in five patients dying of cancer[8,9]. In patients with isolated cardiac localization, allogenic heart transplantation may be meaningful as a potential curative approach[6,10]. For early diagnosis, a broad echocardiographic screening in patients with even mild cardiac symptoms, like palpitations as in this case, might be reasonable.
REFERENCES


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