LV Myxoma on Real-time 3DTTE in a Young Patient with Dilated Cardiomyopathy and Apical Thrombus

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Myxoma, 2DTTE: Two-dimensional Transthoracic Echocardiography, RT-3DTTE: Three-dimensional Transthoracic Echocardiography, LV: Left Ventricle.

Introduction
LV myxoma is extremely rare disease, surgical resection of myxoma always indicated because of high risk embolic events. LV myxoma may be occasionally detected or by the symptoms caused by embolic events or valvular orifice obstructing. Previously 2DTTE was used to make diagnosis prior to respect the tumor. Currently, 3DTTE overcomes the limitations of 2DTTE in description of shape, size, pedicle, location, relationship with around structures, especially the surface aspect of myxoma.

Case Report
A young man patient of 31 years presented at my hospital by heart failure symptoms for one month, including dyspnea on exertion, orthopnea. Physical examination detected an early diastolic gallop sound. No past history of tabac, alcohol consumption, systemic hypertension or congenital cardiac defect had been reported.

He was evaluated immediately by a chest X ray that demonstrated a cardiomegaly with enlargement of the left inferior cardiac arch.

His ECG showed sinus tachycardia and LV hypertrophy with Sokolow-Lyon Index=65mm.

The transthoracic echocardiography initially revealed enlarged LV with LVDD =71mm, severe reduced systolic function of left ventricle: EF=20%, the global contraction of LV diminished severely on par sternal views as well as all apical views.

The two-dimensional echocardiography also recorded an apical thrombus of 13mm x 9mm in diameter that presented as an echogenic mass attached to the apical endocardium.

The Real-time three-dimensional TTE then have been performed with Live 3D, Full Volume and especially 3D Zoom mode that provided 3D image of the hyperechogenic mass about shape, size, mobility, surface, location very clearly. Based on characteristic findings of the echogenic mass as highly mobile, pandiculated, tumor with excrescence aspect, the first diagnosis is LV myxoma.
Figure 1: Cardiomegaly with enlarged the left inferior arch (dilated LV).

Figure 2: ECG showing sinus tachycardia and LV hypertrophy with Sokolow-Lyon Index=65mm.

Figure 3: Transthoracic TM mode demonstrating dilated LV with severely reduced global systolic function.
Figure 4: Apical longitudinal view showed: a- an apical thrombus 13x 9mm, b-an hyperechogenic mass 27x14 mm with narrow stalk adjacent to posterior IVS.

Figure 5: Irregular surface and larged stalk hyperechogenic mass attached to IVS illustrated by Two-dimensional Echocardiography.

Figure 6: 3DTTE apical 4 C view showed an echogenic mass next to the apex.

Figure 7: 2D TTE X-plane mode detected the hyperdense echogenic mass inserted to the posterior IVS near the apex.

Figure 8: Real-time 3DTTE Live mode viewed from apex to mitral valve, the 3D image visualized pedunculated echogenic mass attached next to the apical IVS.

Figure 9: Real-time 3DTTE visualized the echogenic mass that having the narrow pedicle, villious excrescences with high risk of embolic event in compare with polypoid mass.
Primary cardiac tumors are rare, present in less than 0.2% of unselected autopsy series and associated with high frequency of embolic events. Two most common varieties of these tumors are myxoma and papillary fibroelastoma. Approximately 75% of all cardiac tumors are benign and 50% of them are myxoma. It is estimated that 30-40% of all myxoma will embolize [4].

The majority of myxomas are in the left atrium (75%), then right atrium (20%) and 2.5% reported for myxoma in left ventricle [4]. Large, mobile, villous excrescences myxomas are more likely embolize than those with polypoid aspect. The risk of embolic events of our patient surely are very high, but the myxoma cannot resected now. We avoid performing TEE and continue follow up patient with medical treatment of heart failure regarding the severe heart failure of the patient.

Wei Qin et al. [5] reported a LV myxoma in an asymptomatic patient 60 years male, the tumor located at base of anterolateral papillary muscle detected by 2D TTE, the tumor was resected successfully and no recurrence noted.

Solmaz et al. [2] reported a large LV myxoma in a 27 years woman presenting with epigastric pain and weigh loss. This tumor fully filled the LV and compromised the flow of the aortic and mitral valve. The patient was performed 2DTTE then 2DTEE, her tumor originating from LV lateral wall. Surgical procedure was performed and postoperative was without complication.

Hani M. Mahmoud et al. [3] reported a rare case of big left ventricular myxoma presenting with a cerebral vascular stroke in a 35 y.o male patient. The tumor was detected by routine 2DTTE, its attached to the inferior septum and has been bulging through the aortic valve. Surgical resection done; his neurological status improved after surgery.

Enrico Natale et al. reported a case of LV myxoma originating from the interventricular septum and obstructing the left ventricular outflow tract in a 60 years woman. The big myxoma was demonstrated by 2DTTE, its attached to the apical portion of the IVS, extending into the left outflow tract and prolapsed through the aortic valve. The myxoma was resected and postoperative was uneventful.

Occasionally, LV myxoma in my patient was diagnosed unexpectedly because with the the size of myxoma and its location, this tumor did not protrude the mitral orifice like large LA myxoma, but as presented above, this myxoma likely results in cardiac embolic events.

The role of 3D TTE in this case report is the noninvasive imaging method can detected the myxomas regarding size, location, and especially the 3D TEE can visualize the high risk findings related to embolic events better the 2DTTE.

**Conclusion**

Left ventricle myxoma is rarely seen, patients are commonly are asymptomatic and the tumor may be detected by routine echocardiography.

The patient may be presented at hospital by the cardiac embolic events or hemodynamically compromised symptoms caused by protruding or obstructing the LVOT, aortic and mitral orifices, particularly in case of big LV myxoma.

In majority of cases, LV myxomas are pedunculated and attached to the interventricular septum. The big LV myxomas originating from the base of IVS can obstruct the valvular orifices. The small and apical IVS originating myxoma are usually asymptomatic.

The 3DTTE with advances in image processing and display has increased the availability and utility in detecting LV myxoma and providing the important information’s before surgery as well as predicting the risk of embolic events, especially RT-3DTTE records the mobility degree ,the pedicle size and shaped, the location and the villous excrescences of tumor better the 2DTTE used only.

**References**