Have you seen a case like this?

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- A tall statured 17 Y old previously healthy boy was transferred to our hospital from another facility on ventilator with diagnosis of Septic shock, ARDS and presumed infective endocarditis. bed side echo there, revealed a mass attached to TV and MVP with severe MR.
- His family denied history of any high-risk behavior for sexually transmitted diseases, alcohol abuse, intravenous drug abuse, recent dental procedures, persisting skin infections, congenital heart disease, or rheumatic fever.
- At our institution, the physical exam was remarkable for pan-systolic murmur over the apex, bilateral basal to mid-zone lung crackles. His physique was remarkable for disproportionate ration of the lower extremities to torso and very large upper extremities span, he has a high arched palate and the thumb wrist test was positive all suggested marfanoid habitus. His initial laboratory data showed significant leukocytosis and elevated inflammatory markers and microscopic hematuria. Blood, respiratory and urine cultures were negative. Chest imaging showed pulmonary edema like picture/ARDS. TTE and TEE revealed large sessile cauliflower like mass attached to ventricular side of TV anulus and septal leaflet with highly mobile sphere like mass attached to its tip without hemodynamic compromise, Flail anterior MV leaflets with ruptured chordi at A2,A3 scallops and severe MR, Aortic valve showed retraction of RCC with triangular gape causing severe AR
- The consensus was for surgery to remove the mass and send for C/S and histopathology, repair/replacement of valvular lesions.

- Vegetations were removed and sent for culture and histopathology. All culture specimens were negative, Aortic valve specimen revealed No evidence of IE, only myxoid changes (C/W connective tissue diseases). Both aortic and mitral valves were replaced by mechanical valves.

We did a literature review about native TV Endocarditis with atypical vegetation at ventricular side and if there any relation to connective tissue disorder like Marfan syndrome. We found that, Isolated tricuspid (TV) endocarditis accounts for 5%-10% of cases of infective endocarditis (IE) and is uncommon in an immunocompetent adult in absence of risk factors or CHD. Persistent fever associated with pulmonary events, anemia, and microscopic hematuria is known as 'tricuspid syndrome', and should alert for TVE. Early Echo is recommended in such patients. Sometimes, atypical presentation of vegetations at ventricular side of TV may occur in some Patient with VSD and L-> R shunt which Encroach on the Papillary Muscle and Right Ventricle Cavity. Echocardiography is the mainstay of assessment of Marfan's syndrome which may include aortic valve with Annulaoaortic ectasia, especially with dilatation of aortic root, is found in 60% to 80% of adult cases which can cause severe AR or may progress to aortic root dissection. Also Mitral valve may suffer from MVP which is less benign than the common type of MVP identified in the general population. Flail leaflet is an independent predictor of progression of MR and MV-related clinical events.

Back to our case, we found no single case report in the literature with combination of those rare findings.