Figure 1. Static image from video showing jugular venous distention to earlobes with cannon V waves.

A 66-year-old man experienced recurrent ascites of unknown etiology over six months. He had previously undergone a renal transplant secondary to complications of diabetes and hypertension and had known severe coronary artery disease. His most recent paracentesis revealed an albumin 1.6 g/dL (serum albumin 2.1) and a total protein of 3.8 g/dL. His adenosine deaminase was 11.6 U/L (normal <7.6 U/L), but repeated bacterial and mycobacterial ascites cultures were negative, as were a carcinoembryonic antigen assay and ascites cytology. Computed tomography of the abdomen showed findings consistent with cirrhosis, but an extensive workup for common causes of cirrhosis was negative.

Physical exam showed jugular venous distention with prominent V waves and a holosystolic murmur at the left lower sternal border (Figure 1). Echocardiography showed a dilated right ventricle, moderate pulmonary and tricuspid regurgitation and an estimated right ventricular systolic pressure of 87 mm Hg. Cardiac catheterization confirmed the presence of an elevated right ventricular pressure of 72/10 (22) mm Hg, an elevated pulmonary artery pressure of 75/27 (45) mm Hg and a left ventricular ejection fraction of 20-25%. The right atrial pressure was 20 and the pulmonary artery occlusion pressure was 22 mmHg. A diagnosis of pulmonary hypertension secondary to left ventricular heart disease (type 2 pulmonary hypertension) with congestive hepatopathy and cardiac ascites was made.
The patient’s physical examination provided an important clue to the etiology of the ascites – cardiac ascites is thought to be due to chronic venous congestion of the liver due to transmission of high central venous pressures. Tricuspid regurgitation can be associated with severe hepatic congestion because of retrograde transmission of right ventricular pressure directly into the hepatic veins. In some patients (although not in this patient), careful examination will reveal that the liver in such patients is palpably pulsatile.

Cardiac ascites is typically characterized by a serum albumin gradient (SAAG) \( \geq 1.1 \text{ g/dL} \) (indicative of portal hypertension) and ascites protein level of \( \geq 2.5 \text{ g/dL} \) (1). We cannot fully explain why this patient’s SAAG was low. A complete workup for infectious and oncological etiologies of low SAAG ascites was negative. It has been noted that in patients with known cirrhosis (as in this patient), the finding of a low SAAG has a low specificity for infectious and oncological etiologies of ascites (2). Serositis which can sometimes manifest as ascites can also be a complication of tacrolimus which the patient was receiving s/p renal transplant. It’s possible that tacrolimus might have changed the nature of the ascites fluid in this patient but this is conjectural.

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References