2012

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Recommended Citation
Shah, MD, Paurush; Shah, MD, Gunjan; Chandra, MD, Avinash; Lee, MSIV, Lawrence; and Marhefka, MD, Gregary (2012) “Non-ST Elevation MI as a Unique Presentation of Angioimmunoblastic T-cell Lymphoma,” The Medicine Forum: Vol. 13, Article 6. Available at: http://jdc.jefferson.edu/tmf/vol13/iss1/6
Non-ST Elevation MI as a Unique Presentation of Angioimmunoblastic T-cell Lymphoma

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Case

A 61-year-old Chinese female with a history of hypertension, hyperlipidemia, asthma, and gastroesophageal reflux disease presented with four days of chest pressure that radiated to her left arm and jaw. On exam, her vital signs were within normal limits and cardiac and pulmonary exams were unremarkable. Her initial electrocardiogram (ECG) demonstrated ischemic ST segment depressions in leads II, III, and aVF; her first troponin I was elevated at 2.3 ng/mL (normal <0.05 ng/mL) and peaked at 6.8 ng/mL. She was given sublingual nitroglycerin and metoprolol, which controlled her symptoms, and she was started on a heparin infusion to prevent further ischemia. The patient’s medications included: amlodipine was notable for the following daily medications: amlodipine 5 mg, olmesartan 20mg, atorvastatin 20mg, esomeprazole 20mg, montelukast 10mg, and mometasone 110mcg twice daily. Her family history was significant for a brother who had a coronary stent placed at age 57. She denied drug, tobacco, or alcohol use. She was non-English speaking, and immigrated to the United States from China five years ago.

The day after she presented to the hospital, she underwent cardiac catheterization, which revealed diffuse coronary artery disease with the following arterial stenoses: 50% distal left anterior descending (LAD) and 50% mid right coronary artery (RCA) with a 100% distal vessel supplied by extensive left to right collaterals. No interventions were performed. An echocardiogram (echo) was performed which noted equivocal apical septal hypokinesis. Of note, there were circular echodensities with a central echo-free space in the left and right atrioventricular grooves of unclear significance. Cardiac gated computed tomography angiography (CTA) was recommended as an outpatient. The patient was medically managed for a non-ST elevation MI (NSTEMI) and discharged home. Further evaluation of the atrioventricular densities seen on echocardiogram was to be done as an outpatient.

Approximately three weeks later, the patient presented again to the emergency department with recurrent angina, which awakened her from sleep. The ambulance ECG was notable for dynamic ST segment depression, though her troponin was initially within normal limits. The patient was medically stabilized and cardiac catheterization was urgently performed, which again revealed marked progression of coronary stenoses: 90% mid and 50% distal LAD, 90% first diagonal, 90% distal left circumflex, 90% first obtuse marginal, 90% third obtuse marginal, 70% mid RCA, and subtotalted distal right coronary now with right to left collaterals. A total of 300 mcg of intracoronary nitroglycerin was infused with no change in the stenoses. No other interventions were performed and the patient was aggressively treated for coronary vasospasm with diltiazem and isosorbide mononitrate. Her metoprolol was discontinued. Two days later, while medically stable, she underwent a repeat cardiac catheterization, which now revealed resolution of the coronary vasospasm with the same stenoses as her original cardiac catheterization three weeks prior. A rheumatologic work-up was initiated given the unusual atrioventricular densities seen on echocardiogram with vasospasm, and this may have suggested a possible vasculitis. The patient was discharged home again with close follow-up while the rheumatology labs were pending.

Three weeks later, the patient presented again with early morning angina and was treated medically for coronary vasospasm, as well as a recurrent NSTEMI. The patient was stabilized and a cardiac gated CTA was performed, which revealed sleeve-like soft tissue densities encasing the right and left circumflex coronary arteries as well as the proximal portion of the LAD (Figure 1). The prior rheumatologic work-up revealed only a modestly elevated erythrocyte sedimentation rate (ESR). Rheumatology was consulted and they suggested biopsy for tissue diagnosis though one was not performed at that time. A computed tomography (CT) of the chest, abdomen, and pelvis was done revealing mediastinal and para-aortic lymph nodes. A cardiac gated magnetic resonance imaging (MRI) was performed which further characterized the known atrioventricular soft tissue densities (Figure 2). On hospital day 9, the patient developed late evening angina with recurrent ECG ST depressions, and despite sublingual nitroglycerin and eventually an intravenous infusion, the patient became rapidly hypotensive, lost her pulse and suffered a pulseless electrical activity cardiac arrest with failure to respond to resuscitation.

The family agreed to an autopsy which revealed angioimmunoblastic T-cell lymphoma (AITL) with involvement of the coronary epicardium and nodular perivascular infiltration of the right mainstem, left circumflex, and left anterior descending coronary arteries, kidneys, and spleen, as well as mediastinal, omental, mesenteric and para-aortic lymph nodes. (Figure 3 A-D) Pathology revealed the classic picture for AITL, polymorphic infiltrate of lymphocytes, plasma cells, and eosinophils (Figure 4B). Other findings included proliferation of venules and follicular dendritic cells, monoclonal/oligoclonal T-cells present (CD3+/ CD4+), and sometimes oligoclonal or monoclonal B-cell populations attributed to B-cells infected with Epstein–Barr virus (EBV) (Figure 4 A-C).

Discussion

Recurrent NSTEMIs secondary to coronary vasospasm with echodense masses seen in the atrioventricular grooves is a unique presentation with a varied differential including malignancies and rheumatic diseases. More frequent cardiac imaging and biochemical laboratory testing have increased the incidence and prevalence of cardiovascular diseases found in patients with rheumatic disease.1 Systemic lupus erythematosus,
Takayasu disease, rheumatoid arthritis, polyarteritis nodosa, Churg-Strauss syndrome, Behçet’s disease, sarcoidosis, and giant cell arteritis are some of the many reported causes of coronary artery vasculitis. Additionally, a case of lymphoplasmacytic arteritis of the coronary and internal mammary arteries with no systemic disease has also been documented. Myocardial ischemia and infarction without serological evidence of inflammatory disease, as in this patient, unfortunately do not have established management strategies since revascularization in such cases is often unsuccessful.

Tumors with cardiac involvement may affect all three layers of the heart. Metastatic neoplasms commonly involve the epicardium or pericardium, while the less common primary cardiac malignancies often involve the myocardium or cardiac chambers.

The most common primary benign masses of the heart include myxoma, rhabdomyoma, fibroma, hemangioma, and atrioventricular nodal, while the most common malignant neoplasms include sarcomas (angio, unclassified, fibrous histiocytoma, and osteo) and lymphomas. Cardiac involvement from melanoma, germ cell, leukemia, lymphoma, and lung cancers occur in up to 45% of patients with cardiac metastasis, though these may go unrecognized until autopsy. Direct extension, hematogenous spread, retrograde lymphatic extension, or transvenous extension are the accepted mechanisms of metastasis. Symptoms are based on the tumor’s location and often include shortness of breath or chest pain, but most commonly patients present with a pericardial effusion with tamponade. A preliminary diagnosis can be made by echocardiography; CT and MRI are used for further classification, and biopsy is required for confirmation. Treatment of the primary malignancy, whether surgical excision or medical management, depends upon the tumor classification.

AITL, though rare, is one of the more commonly occurring peripheral T-cell lymphomas and accounts for 2-5% of all non-Hodgkin’s lymphomas. It arises from peripheral CD4 positive T-cells thought to be a subset of follicular helper T-cells. Patients tend to be diagnosed after a subacute illness in the sixth or seventh decade with a slight male predominance. Presenting signs and symptoms include asymptomatic generalized lymphadenopathy, fevers, night sweats, arthralgias, and weight loss. Extra-nodal involvement is common and usually includes hepatomegaly, splenomegaly, pleural and pericardial effusions, and various skin rashes. Laboratory abnormalities include an elevated serum lactate dehydrogenase level, a polyclonal hypergammaglobulinemia, and Coombs positive autoimmune hemolytic anemia. Diagnosis is made by excisinal tissue biopsy. Lymph node histology will show an effaced nodal architecture, prominent arborizing high endothelial venules, and
a polymorphous infiltrate, which includes reactive lymphocytes, immunoblasts, plasma cells, and atypical lymphocytes positive for CD3 and CD4. Therapeutic options have not been standardized, but responses have been reported in those receiving single agent corticosteroids, low-dose methotrexate, fludarabine, lenalidomide, and others. Combination chemotherapy, including regimens such as vincristine, doxorubicin, and prednisone with or without cyclophosphamide, has failed to increase survival rates to greater than 30%.

Upon performing a literature review for AITL, we found no other cases in which the presenting symptom was a myocardial infarction. For extranodal manifestations of AITL, there is a case of syncope due to sinoatrial node disease. The authors concluded that the syncope was associated with sinus arrest due to ischemic damage and fibrosis of the sinoatrial node secondary to AITL. Our patient’s diagnosis was especially challenging given that her workup only demonstrated non-palpable lymphadenopathy and echodense masses in the atrioventricular grooves. These masses caused extrinsic compression of the coronary vasculature resulting in myocardial ischemia, repeated infarction, and ultimately to her death.

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