Left Main Spontaneous Coronary Artery Dissection: The Complexity of Management Considerations

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Abstract

Spontaneous coronary artery dissection (SCAD) is a rare, complex clinical entity that is often a diagnostic dilemma. Additionally, SCAD is often encased in uncertainty in terms of the optimal therapeutic approach. We present a challenging case of left main coronary artery dissection outlining how clinical presentation; angiographic features of the dissection, and the degree of obstruction to flow delineate the best clinical approach.

Abbreviations: ED: Emergency Department; CABP: Coronary Artery Bypass; LAD: Left Anterior Descending; LM: Left Main; RCA: Right Coronary; CF: Artery Circumflex; IABP: Intra-Aortic Balloon Pump; LIMA: Left Internal Mammary Artery; SV: Saphenous Vein; SCAD: Spontaneous Coronary Artery Dissection; PCI: Percutaneous Coronary Intervention; FMD: Fibro Muscular Dysplasia

Case Presentation

41-year old white female presented to the Emergency Department (ED) with sudden onset, substernal chest pressure, which she characterized as 10/10, radiating to her upper back. She experienced associated nausea and diaphoresis. She was vacuuming the living room at her home when symptoms developed. The patient’s only risk factor was cigarette smoking 1ppd. Her initial ECG showed sinus rhythm with ST-segment elevation in lead aVR, with reciprocal ST-segment depression in the anterior and inferior leads (Figure 1).

She was given 2 sublingual nitroglycerin tablets in the ED with resolution of her symptoms. Her ST segment ECG changes normalized in the ED, but sinus tachycardia persisted. Her point of care troponin was elevated at 23.8ng/mL (Normal range: 0.0-0.23ng/mL). Her other lab tests, including her CBC and CMP, were normal. She was hemodynamically stable, with blood pressure of 140/80, and her cardiac physical examination was normal. A stat echocardiogram revealed preserved left ventricular [LV] function without regional wall motion or valvular abnormalities. Doppler data revealed elevated LV filling pressure. She was taken

Figure 1: Sinus rhythm with ST-segment elevation in lead aVR, with reciprocal ST-segment depression in the anterior and inferior leads.
Post catheterization the patient was symptom free and hemodynamically stable. The clinical team’s decision was to observe the patient. The patient remained stable and a decision was made to repeat the cardiac catheterization three days after initial presentation. The angiogram revealed no change in the LM dissection, and there was persistent 50% obstruction with TIMI 2 flow. The interventional cardiologist decided the LM stenting was high risk and cardiovascular surgery was consulted. The next day while waiting for Coronary Artery Bypass (CABG) she became hemodynamically unstable and developed pulmonary edema. The ECG revealed ST depression inferiorly and laterally. She had emergent Intra-Aortic Balloon Pump (IABP) placement and CABG with a Left Internal Mammary Artery (LIMA) to the LAD, and Saphenous Vein (SV) grafts to the ramus artery and left CF. Post operatively she experienced bleeding and returned to the operating room for exploration where a vein was clipped at a leak site.

Post op echocardiogram revealed depressed ejection fraction (LVEF of 25%). There was diffuse hypokinesis of all regional segments of the LV and a large apical filling defect consistent with LV apical thrombus. She remained in the hospital for 14 days post op. The patient returned to the hospital in two weeks with recurrent chest pain. Contrast echocardiography revealed resolution of the apical clot. Repeat angiography revealed the grafts were occluded. The patient was referred to another medical center for LM stenting.

Discussion

Spontaneous Coronary Artery Dissection (SCAD) is a rare condition identified in a myriad of case reports and retrospective studies. Most publications cite its incidence to be approximately 0.2%-1.1% [1]. It is relatively well known, at this point in time, as a rare condition that mostly affects women in their third through fourth decades of life [1-4]. The incidence of SCAD seems to be higher in Caucasian women [3,5]. Several studies have reported increased frequency in peripartum women [4]. Despite being relatively well characterized from an epidemiological standpoint, several questions remain regarding the disease processes’ pathophysiology, risk factors, and optimal definitive management.

Some investigators have suggested SCAD’s pathophysiology is related to hematoma formation in the vessel wall. This can occur in the wall of the media, or between the media and adventitia [6,7]. Subsequent expansion of this arterial space can compromise the true lumen and eventually lead to myocardial ischemia. However, many other mechanisms have been implicated ranging from intimal tears to alterations in collagen synthesis [8]. The exact pathophysiology remains an enigma.

The most common association identified in the literature is the association between SCAD and peripartum patients [2-4]. Studies have identified an incidence of SCAD in peripartum patients ranging from 30-48% [2,4]. A lack of traditional risk factors for Coronary Artery Disease (CAD) has also been shown to be common in patients diagnosed with CAD [1-4]. Of traditional CAD risk factors, smoking and hypertension were most commonly identified [4,5].

Saw et al. identified Fibromuscular Dysplasia (FMD) as a possible predisposing condition to SCAD. In their study, they found that 86% of their cohort had FMD in greater than or equal to one non-coronary territory. Non-coronary vascular territory was defined as renal, iliac or cerebrovascular. Even more, 42% had FMD in greater than two non-coronary vascular territories, suggesting that perhaps clinicians should have increased suspicion of SCAD in these patients.

Identifying unique presenting clinical features and symptomology of patient’s eventually diagnosed with SCAD remains of the utmost importance for providers who first encounter such patients. Unfortunately, an overwhelming majority of this patient population presents with symptoms typical of Acute Coronary Syndrome (chest pain, arm pain, SOB). They also typically present with elevated cardiac markers [1-5]. This serves to be particularly challenging as it limits clinicians’ ability to make the diagnosis clinically.

The diagnosis of SCAD can be made in a number of objective ways. A few methods include coronary angiography, CT angiography, optical coherence tomography, and Intravascular Ultrasound (IVUS) [6]. Once diagnosed, SCAD can be further classified into Types I-III. Classification is based on extent of coronary artery involvement, severity of artery stenosis, and presence of intramural hematoma [5,6].

Numerous of studies have sought to determine optimal management for SCAD, however, the consensus remains that there is no definitive superior treatment. At the most basic level, treatment approach varies between medical management and invasive therapies, specifically Percutaneous Coronary Intervention (PCI) and CABG. Several studies suggest that a conservative approach with close follow-up is superior for stable, non-obstructive lesions [9-11]. There is also evidence to suggest that a more aggressive approach (revascularization) is favorable in the setting of ischemia [11]. Ultimately, each of these studies reaches the same conclusion: more data is needed to confidently recommend a particular treatment modality. Overall, the LAD is affected in 75% of cases, the RCA in 20% of cases, CF in 4% of cases, and the left main coronary artery in <1% of cases [12-16].

Revascularization for SCAD is warranted in those patients who present with ongoing ischemia refractory to medical treatment. Cases of spontaneous left main dissections reported in the literature have mainly been treated with CABG, [2-7] especially those with RCA involvement 15 and triple-vessel CAD.16 Successful revascularization with percutaneous stenting has also been described. [17-19]. In cases of severe heart failure, bridging...
to orthotopic heart transplantation with a Left Ventricular Assist Device (LVAD) may be the only option.

Conclusion

This case represents a classic example of the clinical quandaries associated with LM SCAD. This patient presented with hemodynamic instability, a significant LM dissection, a 50% LM luminal obstruction, diminished coronary flow, and evidence of ischemia as manifest by the biomarkers. Classification based on extent of coronary artery involvement, severity of artery stenosis, and presence of intramural hematoma suggests this represents a class 2-3 dissection. In this circumstance the clinician must determine the best approach to treatment in light of the clinical presentation, the complexity of coronary anatomy, evidence of ischemia, and the skill of the interventional team. In this case the unstable nature of the presentation with ischemia (ECG and biomarkers) and the complexity of the LM dissection suggested the need for “timely” intervention. The normal native coronary arteries raised a concern for graft closure due to competitive flow. In retrospect, the best approach would have been immediate LM stenting by an experienced interventional cardiologist. This case highlights the complex decision making associated with the therapeutic approach to LM SCAD.

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