Coronary artery bridging as an etiology for non-atherosclerotic myocardial infarction: A review of literature and case history

Jeffrey Dwyer*
Clinical Specialist in Cardiology, Department of Medicine, Kaiser Permanent Medical Center Vallejo, California, USA

Abstract
Myocardial infarction may occur without atherosclerotic lesions discernible by coronary angiography. The impact of this phenomenon may not be appreciated by many clinicians largely because of a priori assumptions that patients who present with a history of MI or angina syndrome are victims of atherosclerosis and fit well into mainstream management ranging from acute care to rehabilitation. In the absence of atherosclerosis, myocardial infarction may result from several etiologies by chronic hypoperfusion if the culprit artery has a course within the myocardium rather than a more epicardial course. These vessels, called tunneled arteries, are typically traversed by thick bundles of muscle fibres that comprise a myocardial bridge. They are characterized by chronically abnormal hemodynamics that are prone to exacerbation, leading to anginal symptoms and myocardial infarction. A case history is presented that describes the presentation, medical evaluation and treatment, and rehabilitation of a patient with non-atherosclerotic MI attributed to a myocardial bridge. It is followed by a review of the pathophysiology of this medical problem and efficacy of various treatments. Rehabilitation considerations that apply to patients with a tunneled coronary artery are discussed.

Introduction
Myocardial infarction may occur without atherosclerotic lesions discernible by coronary angiography [1-3]. The impact of this phenomenon may not be appreciated by many clinicians largely because of a priori assumptions that patients who present with a history of MI or angina syndrome are victims of atherosclerosis [1,4,5] and fit well into mainstream management ranging from acute care to rehabilitation. Prominent textbooks [4-6], practice guidelines [7-9], and reviews [2,10-12] give little or no recognition of non-atherosclerotic causes of ischemia heart disease. Even young patients without overt risk factors for atherosclerotic vascular disease are often viewed as members of the predominant cohort [9] or a close variant of it [13]. This occurs despite the high frequency of non-atherosclerotic perfusion disorders seen in people aged 40 years or less who present, in some population centres, with symptoms of ischemia [12]. A lack of appreciation for the distinction between atherosclerotic and non-atherosclerotic ischemia heart disease may diminish the focus of services such as cardiac rehabilitation, limit the clinician’s understanding of the patient’s pathophysiology and risk for future cardiac events, and contribute to a tendency for inadequate evaluation [14] and medical treatment [15,16].

In the absence of atherosclerosis, myocardial infarction may result from several aetiologies including vascular spasm, transient dysrhythmia, substance abuse, hypercoagulability and coronary thrombus formation, and dissection of acquired or congenital vascular anomalies [2,3]. The pathophysiology of these events may be exacerbated by chronic hypoperfusion if the culprit artery has a course within the myocardium rather than a more epicardial course. These vessels, called tunneled arteries [1,17] are typically traversed by thick bundles of muscle fibres that comprise a myocardial bridge [18] (Figure 1). They are characterized by chronically abnormal hemodynamics that are prone to exacerbation [2,14,19] and found in up to 85% of necropsy studies performed when the suspected cause of death was heart disease [18,20]. Among patients who present for coronary angiography after MI or onset of angina syndrome, the incidence of a myocardial bridge is as high as 18% [17], predominantly located in the left anterior descending coronary artery [2,17]. The 5-year survival is reportedly greater than 90% [21,22] but disabling symptoms, recurrent MI, serious dysrhythmia, heart failure, and sudden death are recognized as early sequelae to the discovery of a tunneled coronary artery in a patient with chest symptoms [2,20,23-25].

This article offers a case history that describes the presentation, medical evaluation and treatment, and rehabilitation of a patient with non-atherosclerotic MI attributed to a myocardial bridge. It is followed by a review of the pathophysiology of this medical problem and efficacy of various treatments. Rehabilitation considerations that apply to patients with a tunneled coronary artery are discussed. Several case histories are available [15,16,26-30] but none of them include information about the specific rehabilitation of these fascinating patients.

Case history
KD is a 45-year-old female who experienced sudden onset, sharp, substernal chest pain while teaching at a middle school. The pain radiated to both shoulders and did not abate with rest. Prior to this event, the patient had been active without chest symptoms during
strength training, jogging, running, and skiing. KD drove herself to a medical clinic where she was not seen by a physician for 45 minutes due to her failure to conform to the usual stereotype for a patient with ischemic heart disease. An ECG revealed inverted T waves, infero-laterally, with abnormal R wave progression. She was admitted and ruled in for a non-ST elevation MI.

Aspirin, metoprolol 50 mg BID, and lisinopril 10 mg QD were started but intermittent chest pain persisted. At one time the pain was rated 6/10 and associated with systolic BP of 80 mm Hg and bradycardia. The beta-blocker and ACA inhibitor were held and isosorbide dinitrate 10 mg BID was started. An echocardiogram performed one day post-MI revealed a large area of antero-apical hypokinesis with ejection fraction of 50%. No mural clot was found but the patient was considered at risk for this post-MI event. Coumadin was started, and aspirin discontinued.

Risk factor assessment revealed that the patient had smoked 0.5 packs-per-day for ten years but quit 17 years prior to her MI. None of the traditional CAD risk factors were present. This very active woman had an HDL-CHOL of 81 mg/dl, TOT-CHOL of 207 mg/dl, and BMI of 22.5.

Post-MI day 1, an angiogram revealed an ejection fraction of 50% with severe hypokinesis of the apical anterior wall and distal third of the inferior wall with aneurysm dilatation of the apex. The left main coronary artery (LM), left circumflex (LCX), and right coronary (RCA) arteries were angiographically normal. The anterior descending coronary artery (LAD) demonstrated a long segment bridged by myocardium. This segment rounded the apex of the heart and demonstrated complete obliteration of the vessel during systole. No intervention was performed.

Fourteen days post-NSTEMI, the patient was evaluated in the cardiac rehabilitation centre. She reported frequent episodes of chest pain characterized as “pinching” sensations with light activity despite strict compliance with medical therapy. Her exercise prescription included a target HR of 84-96 bpm (14-16 beats/10 second count), equivalent to 20% of the HR reserve. This target HR regulated exercise at 3.5 METs, a pace that maintained the patient below her symptom threshold. KD continued cardiac rehabilitation twice each week for 8 weeks. During that time, the severity of her chest symptoms declined while the symptom-threshold increased to 7 METs with a HR of 118 bpm (46% HR reserve).

Eight weeks post-MI, the patient was advised that she must remain on coumadin indefinitely and that no additional medication, intervention, or follow-up echocardiogram was warranted despite limiting chest symptoms that occurred in bursts of activity, such as climbing stairs, which failed to diminish with 1 to 2 nitro-glycerine tablets. A second medical opinion was obtained, and a follow-up echocardiogram was obtained. Anterior and inferior wall segments were now mildly hypokinetic with nearly complete resolution of the aneurysm dilatation of the apex. No thrombus was seen. Coumadin was discontinued.

**Discussion**

Myocardial bridges are congenital [18,33,34] yet symptoms of ischemia are rare in patients younger than 35 years [17,19,35]. Investigations employing intravascular ultrasound and other modalities have explored this dichotomy and described the mechanisms of ischemia and factors that may lead to serious cardiac events. Features of a tunneled artery that may be viewed as risk factors for symptomatic ischemia or myocardial infarction include age of the patient, length of the muscle bridge overlying the artery, degree of systolic compression of the vessel, number of segments or arteries tunnelled, high heart rate, degree of general myocardial hypertrophy, and depth of the tunnelled vessel [2]. Recent clinical investigations have identified successful therapeutic options and the risk of secondary events [35].

**Angiography**

Among patients who undergo coronary angiography after MI or ischemia, the incidence of a single myocardial bridge ranges from 1 to 18% [36-39]. This is in sharp contrast to several reports of an incidence of 58-86% detected on autopsy in persons suspected of dying from a cardiovascular disorder [17,18,35-41]. This discrepancy may result from variations in muscle fibre orientation of the myocardial bridge, bridge length, and thickness [18], degree of luminal area reduction during systole and early diastole [15,18,42,43], and geometry of the segment during maximal compression [14,17]. Several investigators suggest that angiographic technique, including analytical and observational skills of the operator, contributes to success in identifying hemodynamically significant myocardial bridges [15,17,44]. Standard angiography may result in poorly visualized tunnelled arteries or incorrect measures of their characteristics and propensity for ischemia. When provocative drugs such as nitro-glycerine [38,45], glycercyl dinitrate [46], dobutamine [44,47], and epinephrine [17] are administrated, myocardia bridges are more readily detected.

Using quantitative angiography, several studies involving a total of 80 patients indicates systolic reduction of the diameter of the bridged LAD artery by 71-83% [17,42-44]. Some patients demonstrated greater than 90% luminal diameter reduction with atrial pacing at 134 bpm [19]. In early diastole, the luminal diameter remained reduced by 34-41% [19,42-44]. Noting the limitations of angiography, Ge et al. [14] used intravascular ultrasound (IVUS) to assess changes in luminal area throughout the cardiac cycle. In the normal vessel, proximal to
the myocardial bridge, luminal area varies 9.0 ± 7% from distal to systole. The greatest gain occurred in mid-systole coincidental with peak pressure in the aortic root. In contrast, tunneled arteries have a variation of 40 ± 7%, reaching the minimal lumen area in systole. The loss of luminal area persists in early diastole and becomes more pronounced in tachycardia [19, 43].

Several investigators [14,15,44] suggest that a thorough evaluation of a tunneled artery should include IVUS in addition to angiography. Using IVUS, Ge et al. [14] discovered atherosclerotic plaques in the LAD artery proximal to the myocardial bridge in 12-14 patients. Angiography failed to identify plaques in 8 of these patients. Using standard angiographic techniques and IVUS, Schwartz et al. [19,42] and Klues et al. [43] found no atherosclerotic lesions within bridged segments or normal segments distal to the bridge.

**Hemodynamics**

The phasic loss of luminal dimensions in tunneled arteries described above results in abnormal hemodynamics in both systole and diastole. At peak systole, intra-coronary measurements of maximal pressure reveal a difference of 58.6 ± 43 mm Hg between the point of greatest compression on the tunneled artery and a reference site in the proximal normal vessel coincidental with a sharp decrease in vessel area [43]. This event creates retrograde flow that early investigators labelled the milking effect [18-20]. At the onset of diastole, flow becomes antegrade with a sharp increase in flow rate due to the coincidence of maximal coronary perfusion pressure at a time when the tunneled artery is still compressed by the myocardial bridge [14]. As vessel area increases, flow rate sharply decelerates to a plateau that persists as the myocardial bridge relaxes slowly in the remainder of diastole [19,43]. In contrast, IVUS and Doppler studies focused on the proximal, normal artery indicate that vessel area increases slightly throughout systole as aortic pressure rises [19,42,43,45]. In early diastole, flow accelerates but not sharply as seen in bridged vessels. Throughout mid- and late-diastole, flow declines gradually without displaying a plateau typical of bridged vessels [14].

Coronary flow reserve (CFR), defined as the ratio of mean flow velocity at peak hyperaemia to mean resting flow velocity [35] has been used by several investigators as a sensitive index of the impact of myocardial bridging on coronary hemodynamics [14,19,43] and efficacy of treatment [19,44,46]. Bourassa et al. [35] summarized the CFR and other hemodynamic data reported in five prominent studies. In each study, CFR distal to the bridged artery was significantly impaired, ranging from 2.0 to 2.6 [14,19,42-44]. A normal CFR exceeds 3.0. There is consensus that impaired CFR, attributed primarily to delayed early and mid-diastolic relaxation following severe systolic compression of the vessel, may collectively reduce the threshold for ischemia of the myocardium served by a tunneled artery and explain the occurrence of angina symptoms. Confirmation is provided by Schwartz et al. [19] who measured Doppler flow velocities within tunneled arteries during atrial pacing at 134 bpm. Tachycardia reduced the diastolic period and exacerbated abnormal hemodynamics, eliciting dyspnoea and chest pain. Intra-coronary beta-blocker medication returned all Doppler flow velocities to baseline and resolved nearly all symptoms with ECG changes despite continued pacing. Comparable results were obtained with endovascular stent placement within a bridged artery in patients with severe angina refractory to oral antianginal medical therapy [43,44,46]. Stent placement immediately resolved phasic loss of luminal diameter, systolic and diastolic flow abnormalities, and restored CFR to normal.

**Ischemia and infarction**

Early studies that considered the association between ischemia and compression of a coronary artery by myocardial bridge suggested that these vascular abnormalities are generally innocuous [36,37,48] due to inconsistent correlation between limited angiographic discovery of myocardial bridging and the incidence of clinical events [19]. In the context of abnormal hemodynamics described above, it is now recognized that tunneled arteries can be the primary cause of dysrhythmia [48,50], including atrial fibrillation [51], angina [16,26-29,37], myocardial infarction [24,52-54], and sudden death [24,25,51].

Disparity between the anatomic incidence of bridged arteries and the incidence of clinical events is likely due to an interaction of several fixed and transient factors [15,18,19,26,31,35]. Tunneled segments of coronary arteries greater than 3 cm in length that lie deep in the myocardium are more likely to generate abnormal hemodynamics that lead to ischemia and MI [2,17] than short, shallow tunnels. Moreover, ischemia is more likely to occur when muscle fibres of the over-lying myocardium cross the artery obliquely or helically [18]. This condition may create the greatest degree of vessel distortion during systole and delay diastolic relaxation [55] compared to bridges that cross perpendicularly. Deep bridges with oblique fibres can twist an artery, exacerbating abnormal hemodynamics and creating intimal damage that increases propensity for platelet agregation [56]. Compression of the vessel may be minimal if loose connective and adipose tissue dampens the compressive force of the bridge. If interposed tissue is sparse, vessel compression may be substantial. Aortic outflow tract obstructions may enhance vessel compression due to increased tension in the myocardium that substantially exceeds intracoronary blood pressure. Similar hemodynamics are created if stenosis of the vessel occurs proximal to the bridged segment. Reduced intracoronary pressures distal from the stenotic segment will increase the transmural pressure gradient at the site of tunnelling, favouring vessel compression.

Spasm of a tunneled coronary artery has been cited as a predominant transient factor leading to significant clinical events [2,15,31]. Several investigators implicate heavy cigarette smoking [15,16], extreme alcohol consumption, and substance abuse [58-60] as provocative agents. Many pharmacologic agents may induce spasm of tunneled arteries [17]. Intracoronary injection of dobutamine, epinephrine, or isoproterenol severely exacerbate vessel narrowing in systole and delay of diastolic relaxation [17,47,48]. Similar effects were found with either intracoronary or sublingual nitro-glycerine [38]. The deleterious effect of nitro-glycerine used sublingually by symptomatic patients with bridged arteries was recognized by several investigators [16,27,38].

Several authors [14,38,61] suggest that abrupt, severe tachycardia may create spasm in tunneled arteries. There are no studies, however, that offer hemodynamic or angiographic evidence of spasm in tunneled or normal vessels as a result of spontaneous tachycardia. Schwarz et al. [19] found that atrial pacing at 134 bpm severely exacerbated the hemodynamics in tunneled LAD arteries in 15 patients. Similar results have been found with incremental dobutamine infusion [44].

Coagulation disorders are cited by several authors [15,22,31,57,62] as significant etiological factors in MI not associated with CAD, especially in young people. With one exception, the author found no case histories of patients with MI associated with a tunneled artery that clearly implicated a coagulation disorder. The exception describes a case in which a hyper-coagulable state, resulting from nephrotic syndrome, was associated with MI in a young person without coronary atherosclerosis [63].
Medical therapy

It is generally accepted that negative inotropic agents lower systemic and transmural pressures and reduce the extent of compression of tunneled arteries [23,64]. Schwarz et al. [19] described the effects of intracoronary infusion of esmolol on hemodynamics during atrial pacing at 134 bpm in 12 patients. The beta-blocker reversed the negative effects of tachycardia on average diastolic peak flow velocity within the bridged segment without affecting flow velocity in segments proximal and distal segments. During pacing, average peak flow velocity and diastolic/systolic flow velocity ratio were restored to baseline levels by this medication. Symptoms elicited by the tachycardia were completely resolved in 8 of 9 patients while ECG changes normalized in 5 patients. In contrast to this compelling data, there are no meta-analyses or large studies that describe the long-term clinical efficacy of beta-blocker therapy in symptomatic patients with ischemia or history of MI caused by a bridged coronary artery. Two studies report successful beta-blocker therapy up to one-year post-MI but only three patients were followed [23,27]. Several others have not found beta-blocker to be substantially effective in controlling symptoms of ischemia in post-MI patients with a myocardial bridge [16,26,43-46,61].

If spasm is indicated as an etiologic factor, several authors recommend a calcium channel blocker [16,21,61,65] and an ACE inhibitor or angiotensin receptor inhibitor [21]. Tauth and Sullenberger [28] used diltiazem CD 240 mg/day with success in a 38-year-old man but no long-term follow-up was reported. In contrast, de Winter et al. [16] described a 47-year-old man with chest pain caused by spasm of a bridged artery. He was discharged from the hospital after a negative treadmill test and treated with nifedipine 60 mg daily. Within four weeks he was readmitted with marked ECG indications of ischemia. Similarly, Haager et al. [44] reported failure of anti-angina medical therapy in 9 of 11 patients with angina and a history of MI associated with a bridged LAD artery.

Several authors advise limited use of nitro-glycerine for symptoms of ischemia [16,17,27,38]. Angellini et al. [17] demonstrated severe exacerbation of vessel compression by sublingual or intra-coronary nitro-glycerine. This medicine may enhance contraction of the bridged myocardium, further reducing coronary flow reserve [27,38]. Bashour et al. [27] recommend sublingual nitro-glycerine only for prolonged episodes of severe chest pain associated with ST-segment elevation. The efficiency of long-acting oral nitrates has not been studied. Schwarz et al. [19] suggest that benefits may be derived from reduced preload and suppression of vasospasm.

Invasive treatment

With the advent of intracoronary stents, clinicians theorized that these devices would provide internal stabilization of the lumen of tunneled arteries by counteracting transmural forces and resolving abnormal hemodynamics [46]. Haager et al. [44] stipulated that the stent must be highly flexible or modular, and span at least 90% of the length of the tunneled segment, to provide optimum flow and prevent ischemia.

Several studies have described the clinical course that led to stent implantation in tunneled coronary arteries [15,16,26,44,46]. With one exception, each of these studies included 4 patients or less. Haager et al. [44] studied 11 patients with ischemia or MI attributable to a tunneled vessel. A total of eighteen stents were deployed in the group and the patients followed for two years. Four patients experienced anginal symptoms less than seven weeks after the procedure while a fifth patient was found to have greater than 50% in-stent stenosis without symptoms. Two patients received additional stents and remained symptom-free over the next six months. Two others required surgery with a LIMA graft to the LAD. Several case histories point to high failure rates of stents in tunnelled vessels [14,16,46,66]. Haager et al. [44] suggest that persistent external compression of a tunneled segment might create shear stress that leads to intimal proliferation and in-stent stenosis. Proliferation of the endothelium due to damage incurred during balloon inflation and early mechanical recoil or shrinkage of the stent scaffolding may also contribute to early loss of luminal diameter [26].

In patients with long bridged segments, a surgical option may be preferable to stent deployment [29,44,46,67]. Tio et al. [47] found a 7 cm bridge of the LAD in a 46-year-old survivor of an event they called “sudden cardiac near-death.” Grossly abnormal hemodynamics in this vessel indicated the surgical option. A 5 mm thick muscular bridge was divided without complication. Three weeks later, angiography revealed complete resolution of the bridge and normal hemodynamics. Furnnis et al. [29] also achieved resolution of a LAD bridge by myotomy in a 47-year-old man. Prior to surgery, nuclear imaging demonstrated a stress-induced anterolateral perfusion defect. After ostomy, this defect was not observed, and all symptoms were resolved.

Cleavage of muscle fibres that comprise a bridge requires deep incisions of the ventricular wall and may result in a mural aneurysm [68] or inadvertent opening of the right ventricle [47]. Prendergast et al. [46] suggests that myotomy may be no better than medical therapy because scar tissue development in the myocardium may compress the underlying vessel. Interestingly, Tio et al. [47] observed vessel spasm three weeks post-myotomy. They suggest that this event was the result of inflammation of the vessel wall and/or post-operative scar formation. Pratt et al. [69] recommend coronary artery bypass graft surgery as an alternative to myotomy or endovascular stent placement. They describe two successful cases using the minimally invasive technique. De Winter et al. [16] and Prendergast et al. [46] argue that surgical procedures should be considered only upon failure of an endovascular stent because ischemia from a tunneled artery is usually a single vessel disorder. Unfortunately, the patient described by Prendergast et al. [46] sustained an anterior MI when his stent failed after three years. Haager et al. [44] found that 5 of their 11 patients had in-stent stenosis and abnormal hemodynamics only seven weeks after stent deployment in the LAD. Two of these patients underwent CABG and remained free of symptoms during a two-year period of clinical monitoring.

Cardiac rehabilitation

The cardiac rehabilitation of patient KD was challenging because of basic differences between victims of atherosclerotic ischemia heart disease and those with a myocardial bridge. In contrast to typical patients with anterior MI, KD suffered an infarction at an early age after decades of practicing an active lifestyle that included vigorous aerobic training activities. She possessed none of the traditional CAD risk factors, no intervention was attempted, and no plan was established for rigorous medical follow-up. Furthermore, KD endured an extended period post-MI during which she was limited by anginal symptoms despite medical therapy.

KD was frustrated by the lack of information from her doctors concerning the etiology of her cardiac illness, efficacy of treatment options, prognosis, and the abrupt change from an intensely active lifestyle to one quite limited by symptoms. Discussion of these concerns comprised a significant portion of the education component.
of cardiac rehabilitation. Without CAD risk factors, the usual core elements and goals of traditional patient education course were not relevant. Emphasis was placed on behavior modification with respect to symptom recognition and self-pacing of activities of daily living. Unfortunately, no search was conducted for occult risk factors, or factors associated with non-atherosclerotic MI identified by DaCosta et al. [31].

At 14 days post-MI, patient KD was highly symptomatic at low exercise intensities despite treatment with isosorbide dinitrate 20 mg BID. There are no studies that compare the efficacy of this medicine to calcium channel blockers, or other medications, in this subgroup of MI patients. Sub-lingual nitrates administered for episodes of chest pain have been found to exacerbate the abnormal hemodynamics of tunneled arteries [17,27,38]. However, nitrates may be beneficial in some patients by reducing pre-load and suppression of vasospasm [19]. During KD’s hospitalization and convalescence here was no trial with a calcium channel blocker despite recent literature suggesting this class of medication can be efficacious [28,65].

KD’s angiogram did not identify atherosclerotic stenosis in any coronary artery. Several studies report no angiographic evidence of CAD in bridged vessels associated with symptoms or MI [15,26,28,44,46]. In fact, it has been suggested that tunneled arteries may be protected against the development of CAD [41,70]. Consequently, it is widely held that atherosclerotic disease is not associated in any way with the phenomenon of myocardial bridging. Using IVUS, however, Ge et al. [14] discovered significant atherosclerotic plaques proximal to bridged LAD in eight patients with angiographically normal arteries. It was suggested that the normal, proximal segment may be subjected to wall stress from abnormal flow and shear downstream. Shear stress may injured the vessel wall and lead to intramural plaque formation without loss of lumen [71]. Ge et al. [14] and others [56] suggest that IVUS should be used routinely in patients identified with a myocardial bridge to more fully evaluate the vessel’s status and establish the patient’s risk of future events.

KD’s exercise prescription included an unusually low relative HR based upon her threshold for symptoms in treadmill walking and cycling. Persistent anginal symptoms that improved slowly over several weeks were likely emanating from injured, peri-infarct tissue to which perfusion remained abnormal. At 10 weeks post-MI, KD’s target heart rate was only 46% of HR reserve (~7.0 METS) due to chest pain rated 2-3/10 despite the continued use of isosorbide dinitrate. At 14 weeks post-MI, KD returned to work and resumed an active but very limited lifestyle. Approximately one year later, she was admitted to the hospital with chest pain that began after a session of moderate paced exercise. Her symptoms onset while driving her car and were not relieved with NTG x 2. The second NTG tablet worsened her symptoms. KD ruled out for an MI and performed a treadmill test without indications of ischemia. It was concluded that vessel spasm caused her symptoms but no search for precipitating factors was conducted. Medical therapy was modified to include a low-dose calcium channel blocker with continuation of isosorbide dinitrate.

Summary

Patients who present for cardiac rehabilitation after MI associated with a myocardial bridge may be highly symptomatic. Medical therapy or vascular invention may not offer the same degree of success experienced by patients who have atherosclerotic vascular stenosis as a basis of their MI. Routine evaluations of patients with a myocardial bridge—including angiography—may incompletely describe the patient’s pathophysiology and risk of future cardiac events. Cardiac rehabilitation clinicians may be able to gather data that clearly defines the patient’s functional status and serves as a foundation for medical decision-making processes, particularly about medications.

References

16. DeWinter RJ, Kok WEM, Piek JJ (1998) Coronary atherosclerosis within a myocardial bridge, not a benign condition. Heart 80: 91-93. [Crossref]


29. Furniss SS, Williams DO, McGregor CG (1990) Systolic coronary occlusion due to myocardial bridging-rare cause of ischemia. Int J Cardiol 226: 116-117. [Crossref]


32. Cheitlin, MD, McCallister, HA, de Castro CM (1975) Myocardial infarction without atherosclerosis. JAMA 231: 951-959. [Crossref]

33. Visscher, DW, Miles BL, Walter BF (1983) Tunneled (“bridged”) left anterior descending coronary artery in a newborn without clinical or morphological evidence of myocardial ischemia. Cathet Cardiovasc Diagn 9: 493-496. [Crossref]

34. Von Polacek P, Kralove H (1961) Relation of myocardial bridges and loops on the coronary arteries to coronary occlusions. Am Heart J 61: 44-52. [Crossref]


Copyright: ©2019 Dwyer J. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.