Coronary collateral circulation usually develops as a consequence of recurrent ischemia associated with severe stenosis. In exceptional cases, it can develop with moderate coronary lesions if there is severe recurrent vasospasm. In this situation, the presenting clinical features of vasospastic angina (i.e., effort angina with ST-segment depression) can be identical to those of a severe permanent lesion.

We present a patient who exhibited effort angina and ST-segment depression on treadmill testing. Angiography showed severe right coronary artery stenosis and the development of coronary collateral circulation from the other main artery. After repeated intracoronary bolus injection of nitroglycerin, only a moderate stenosis was apparent and the collateral circulation had disappeared.

**Key words:** Coronary vasospasm. Collateral circulation. Effort angina.

**INTRODUCTION**

Coronary collateral circulation usually appears when stable anterograde coronary blood flow is insufficient. It is frequently found in angiographic studies, generally associated with fixed atherosclerotic obstructions (partial or complete) of coronary arteries. Less common is the development of collateral circulation due to episodes of recurrent ischemia secondary to severe coronary vasospasm in coronary arteries without angiographically significant lesions. We present the case of a patient with a fixed moderate atherosclerotic lesion, associated with severe vasospasm and development of collateral circulation as a consequence of severe recurrent ischemia.

**CLINICAL CASE**

Our patient was a 51-year-old man, a heavy smoker with dyslipidemia, on dietary treatment but without other known cardiovascular risk factors. In recent months he had presented early morning effort angina which become progressively more frequent until it was a daily occurrence. The angina diminished with sublingual nitrates. Electrocardiogram (ECG) and baseline echocardiography results were normal. Clinical and electric treadmill testing results were positive (≤2 mm horizontal ST-segment de-
pression) at 4 minutes of the Bruce protocol. A treatment regimen of aspirin and beta-blockers was established but no improvement in symptoms was observed. Baseline coronary angiography revealed a subocclusive lesion in the mediiodistal right coronary artery (Figure 1), showing the distal vessel through heterocoronary collateral circulation (Figure 2). After bolus injection of nitroglycerin (3 0.2 mg bolus), the lesion was seen to reduce to moderate, its length diminished, and the distal vessel was normal in the anterograde injection (Figure 3). Following a later injection in the left coronary artery, collateral circulation was no longer visible. Subsequently, we started treatment with oral diltiazem and nitrates in transdermal patches, which improved the patient’s symptoms.

DISCUSSION

Coronary collateral circulation is the connection between different segments of the same artery or between different arteries that is established when the original blood flow is insufficient. It develops in response to intermittent myocardial ischemia, protects the myocardium and prevents ischemic events. Myocardial ischemia per se can induce development of collateral circulation, possibly through biochemical signals that provoke the release of angiogenic growth factors. Collateral circulation can help protect the myocardium in patients with coronary artery disease, limits myocardial ischemia during coronary occlusion, and when well-developed can even minimize the infarction zone and predict presence of viable myocardium in patients with a history of previous infarction. This occurs more frequently in patients with severe fixed atherosclerotic coronary artery lesions (occlusions or subocclusions). It is less common to find well-established collateral circulation secondary to episodes of repeated coronary spasm.
even in the absence of significant coronary artery lesions, has been reported.2-5

In vasospastic angina, the classic electrocardiogram pattern is of ST-segment elevation during ischemic episodes.6 Presence of ST-segment depression in the ECG during episodes of angina due to coronary spasm can be put down to subendocardial ischemia caused by partial occlusion of an epicardial coronary artery and the transitory increase of blood flow provided by collateral circulation.7 According to Tada et al.,8 this collateral circulation supplies blood flow thru preexisting vessels to ischemic regions during coronary spasm which prevents transmural ischemia, diminishes the degree of ischemia and is associated with ST-segment depression during episodes of angina. Yamagishi et al.9 confirmed these findings reporting a lower frequency of ST-segment elevation during coronary spasm in patients with established collateral circulation.

Our patient demonstrated effort angina and ST-segment depression on treadmill testing. He had a moderate lesion in the right coronary artery associated with severe vasospasm that caused a subocclusive lesion. Our case highlights the need always to administer intracoronary bolus injection of nitroglycerin in diagnostic coronary angiography to determine the true severity of lesions. Our patient would probably be in “chronic spasm” and ischemia would be seen only on increasing the demand for oxygen during effort. The ST-segment depression in treadmill testing may be due to the lower degree of ischemia thanks to the blood flow contributed by the collateral circulation. Calcium antagonists and/or nitrates are the treatment of choice and were effective for our patient.

REFERENCES