Mesenteric vein thrombosis in a patient heterozygous for factor V Leiden and G20210A prothrombin genotypes.

Paras Karmacharya  
*Reading Hospital*, paraskarmacharya@gmail.com

Madan Raj Aryal  
*Reading Hospital*

Anthony A. Donato  
*Reading Hospital*

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Paras Karmacharya, Madan Raj Aryal, Anthony Donato

Paras Karmacharya, Madan Raj Aryal, Anthony Donato, Department of Internal Medicine, Reading Health System, West Reading, PA 19611, United States

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Correspondence to: Paras Karmacharya, MD, Department of Internal Medicine, 6th avenue and Spruce Street, Reading Health System, West Reading, PA 19611, United States. paraskarmacharya@gmail.com
Telephone: 1-484-6288255 Fax: 1-484-6289003
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Abstract

Mesenteric venous thrombosis (MVT) is a rare but life threatening form of bowel ischemia. It is implicated in 6%-9% of all cases of acute mesenteric ischemia. The proportion of patients with primary (or idiopathic) MVT varies from 0% to 49%, with a decrease in frequency secondary to more recent availability of newer investigations for hypercoagulability. The presence of factor V Leiden (FVL) and prothrombin G20210A mutations (PGM) have been well documented in these cases. However, there have been scarce case reports describing MVT in heterozygotes of both these mutations occurring simultaneously and its implications on long term management.

Core tip: The common presence of two thrombophilic defects increases the thrombotic risk several folds above the risk of a single defect and these tend to occur at an earlier age as seen in our case. Also the risk of recurrent thrombosis is significantly increased among these heterozygotes. Indefinite anticoagulation with oral anticoagulants (goal International Normalized Ratio = 2-3) is recommended for high risk patients like our case with thrombosis at unusual sites (e.g., mesenteric vein), and heterozygosity for both factor V Leiden and prothrombin G20210A mutations. These patients should avoid any hormonal therapy and family members should be screened for underlying prothrombotic condition.


INTRODUCTION

Mesenteric venous thrombosis (MVT) is a rare but life threatening form of bowel ischemia, responsible for 6%-9% of all acute mesenteric ischemia. The presence of factor V Leiden (FVL) and prothrombin G20210A mutations (PGM) have been well documented in these cases. However, there have been scarce case reports describing MVT in heterozygotes of both these mutations occurring simultaneously and its implications on long term management.

CASE REPORT

A 22-year-old Caucasian female presented to the emer-
**DISCUSSION**

MVT is a rare but potentially life threatening cause of mesenteric ischemia with high recurrence rates. It is implicated in 6%-9% of all cases of acute mesenteric ischemia. Predisposing conditions including myeloproliferative disorders, neoplasia, hereditary hemorrhagic telangiectasia, paroxysmal nocturnal hemoglobinuria, inherited thrombophilias, oral contraceptive pill (OCP) use, pancreatitis, recent abdominal surgery or local intraabdominal infections can be identified in most patients. When no underlying etiology is identified, MVT is described as primary or idiopathic. The proportion of patients with primary (or idiopathic) MVT varies from 0% to 49%, with a decrease in frequency secondary to more recent availability of newer investigations for hypercoagulability. Abdominal pain is the most common symptom, especially with acute thrombosis, whereas chronic MVT usually manifests as portal hypertension or diagnosed incidentally by imaging. The increasing use of CT for the investigation of abdominal pain and anticoagulation for the treatment of acute MVT have improved outcomes in these patients. Surgery and bowel resection may occasionally be needed for patients with bowel infarction, perforation, and peritonitis. The management of patients with chronic MVT is aimed at reducing complications of portal hypertension.

The present case is of interest in that acute MVT was the initial presentation in a patient with combined heterozygosity for FVL mutation and the G20210A prothrombin gene variation in the face of oral contraceptive use. The association of each of these mutations with thrombotic disease has been well established. Among Caucasian patients presenting with an initial episode of idiopathic deep venous thrombosis, 12%-20% will be found to be heterozygous for the FVL mutation and 6% heterozygous for the prothrombin G20210A gene variation as compared to 6% and 2% respectively, in asymptomatic Caucasian controls. A recent retrospective study by Amitrano et al noted a high prevalence of thrombophilic genotypes (75%): FVL (25%), prothrombin G20210A gene (25%), and MTHFR prothrombotic defects (50%) in patients with acute mesenteric vein thrombosis. Double heterozygotes of FVL mutation and the prothrombin G20210A gene variation have been shown to be associated with a greater risk of venous thrombosis than either defect alone. Also the age at the first episode of venous thromboembolism in double heterozygotes was significantly younger than those without both gene defects. Thromboembolic disease in double heterozygotes was significantly increased among these heterozygotes of FVL mutation and 6% heterozygous for the prothrombin G20210A gene variation in the face of oral contraceptive use. The association of each of these mutations with thrombotic disease has been well established. Among Caucasian patients presenting with an initial episode of idiopathic deep venous thrombosis, 12%-20% will be found to be heterozygous for the FVL mutation and 6% heterozygous for the prothrombin G20210A gene variation as compared to 6% and 2% respectively, in asymptomatic Caucasian controls.

It seems plausible that in our case, MVT was induced by OCPs on the background of her hematological disorders leading to a hypercoagulable state. The common presence of two thrombophilic defects increases the thrombotic risk several folds above the risk of a single defect and these tend to occur at an earlier age which was also seen in our case. Also the risk of recurrent thrombosis is significantly increased among these heterozygotes. Indefinite anticoagulation with oral anticoagulants (with goal INR = 2-3) is recommended for high risk patients like our case with thrombosis at unusual sites (e.g. mesenteric vein), and heterozygosity for both FVL and PGM. These patients should avoid any hormonal therapy including OCPs due to increased risk of blood clots. It may also be advised to screen the family members for underlying prothrombotic condition, even with a...
first episode of idiopathic venous thrombosis\(^1\),\(^2\).