Portomesenteric venous thrombosis in a postmenopausal female with testosterone implant: a case report

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Abstract

Background: Acute portal vein thrombosis is a rare medical event usually seen in liver disease, but it can also occur due to any inherited or acquired procoagulable state that triggers venous occlusion. Hormonal therapies have been associated with an increased risk of prothrombotic states. This case report documents a portomesenteric venous thrombosis in a postmenopausal woman with testosterone implant for the treatment of hypoactive sexual desire and discusses the importance of identifying hypercoagulable risk factors before initiating hormone replacement therapy. We want to improve the awareness of an unusual medical complication associated with hormone replacement therapy and shed light on how testosterone implants could facilitate a thrombotic event related to other risk factors such as obesity and chronic hypoxic states, as well as the importance of differential diagnosis in the evaluation of postmenopausal women on testosterone replacement therapy presenting with acute abdominal pain.

Case presentation: A 55-year-old obese postmenopausal Hispanic female with medical history of chronic obstructive pulmonary disease presents with intractable abdominal pain, is found to have elevated hemoglobin and hematocrit, and an abdominopelvic computed tomography scan revealing portal and superior mesenteric vein thrombosis. Further evaluation excluded inherited and acquired thrombophilia but revealed elevated testosterone levels. The patient was treated with anticoagulation, which resulted in recanalization of the portal and superior mesenteric veins.

Conclusion: Supraphysiologic levels of testosterone caused by testosterone implants as a treatment of hypoactive sexual desire in postmenopausal women can contribute to thrombotic events in the presence of additional prothrombotic risk factors. Therefore, testosterone therapy should include a thorough risk assessment for prothrombotic states, be tailored to patients’ physiologic testosterone levels, and have close follow-up with testosterone level monitoring.

Keywords: Portal vein thrombosis, Superior mesenteric vein thrombosis, Prothrombotic states, Testosterone implant, Hormone replacement therapy

Background

Portal vein thrombosis (PVT) is a vascular disease that result from mechanical obstruction to the portal vein by a thrombus. Occasionally, the thrombus can extend towards the mesenteric and splenic veins [1–5]. A feared complication is intestinal infarction, which requires prompt surgical exploration [6]. Although uncommon, PVT should be included in the differential diagnosis in the evaluation of postmenopausal women on testosterone replacement therapy (TRT) presenting with acute abdominal pain.

The diagnosis is clinically supported by imaging findings. Ultrasound is the initial examination of choice,
with computed tomography (CT) scan providing additional information [3]. PVT most commonly presents in patients with liver disease, malignancy, and also inherited or acquired prothrombotic states [1–5]. Some medications such as those used for hormonal therapy including testosterone have been associated with increased hypercoagulability [7]. TRT has been used on postmenopausal women to increase libido [8, 9]. A 2019 global consensus position statement by an international expert panel recommended against the use of testosterone formulations, including pellets that may cause supraphysiologic levels, because of the possibility of adverse effects, including increased prothrombotic state [10].

Management of PVT consists of investigating possible causes such as liver disease, malignancy, and prothrombotic states. Additionally, treatment with anticoagulation, thrombolysis, or thrombectomy should start as soon as possible to improve outcomes [2–5]. Prompt diagnosis and treatment lead to a favorable prognosis and reduce the risk of acute and chronic complications such as intestinal infarction and portal hypertension [3, 6, 11].

We report a case that describes portomesenteric venous thrombosis in a postmenopausal female on testosterone therapy to treat hypoactive sexual desire. This case demonstrates that testosterone treatment with supraphysiologic levels can facilitate a thrombotic event in the presence of additional prothrombotic risk factors and highlights the importance of its early detection and treatment.

Case report
A 55-year-old obese postmenopausal Hispanic female, with a body mass index (BMI) of 30.5 kg/m², presented to the emergency department in January 2019 with a 2-day history of diffuse and cramp-like abdominal pain that was worsened by eating. She denied nausea, vomiting, diarrhea, blood in the stool, abdominal trauma, recent surgery, and traveling. Medical history revealed moderate chronic obstructive pulmonary disease (COPD) without a history of recent exacerbation. Medications included Ventolin as needed, Symbicort, Spiriva without any new adjustments, and testosterone pellets implanted 2 months prior for treatment of reduced libido. She denied smoking, alcohol, and drug use. Family history was unremarkable. On physical examination, the patient had oxygen saturation of 95% on ambient air, heart rate 104 beats per minute, blood pressure 135/80 mmHg, and temperature 97.7 °F. Moreover, she exhibited unlabored bilateral breathing with good air entry, and her lungs had bibasilar reduced breath sounds, no chest wall retractions, and no use of accessory muscles.

The abdomen was soft, distended, tender to palpation in the mid-abdomen and right subcostal area, reduced bowel sounds throughout, and no rebound tenderness or guarding. Laboratory studies revealed hemoglobin 17.5 g/dL, hematocrit 52%, leukocytes 9600 per μL, aspartate aminotransferase 55 units per L, alanine aminotransferase 62 units per L, lactic acid 1.4 mmol/L, lipase 98 units per L, a negative fecal occult blood test and basic metabolic panel, urinalysis, and PT/PTT within normal limits. Abdominopelvic CT scan with contrast revealed portal vein thrombosis extending to the mesenteric vein (Fig. 1). Further evaluation excluded known causes of inherited and acquired prothrombotic states. Additionally, testosterone levels were 182 ng/dL. Hematology–oncology suggested a developed prothrombotic state possibly precipitated by TRT, chronic hypoxia, and obesity. Patient treatment consisted of anticoagulation with low-molecular-weight heparin, which gradually improved her abdominal pain. A follow-up abdominal CT scan on day 3 of anticoagulation showed partial portomesenteric vein recanalization (Fig. 2). The patient was later discharged with rivaroxaban for 6 months of anticoagulation therapy and was advised against further HRT. Three months after discharge, she had hemoglobin of 14.5 g/dL and no evidence of prior symptoms or signs of portal hypertension.

Discussion
PVT is an uncommon medical condition with potentially devastating consequences if not recognized early. It has an annual incidence of 0.7–1:100,000 and a lifetime risk in the general population of 1%. Prevalence is higher in males between 45 and 60 years [1]. PVT is defined as the partial or complete thrombotic occlusion of the portal vein, sometimes with extension towards the mesenteric and/or splenic veins. It tends to be more frequent in patients with liver disease but is also seen in patients without liver disease and usually associated with an inherited or acquired prothrombotic state. Presentation of disease can be acute or chronic, with portal hypertension being a complication of chronic disease [1–5]. In more than 90% of cases, acute PVT manifests with progressive abdominal pain and distention. Other commonly accompanying symptoms are fever and ascites. In extreme cases, the patient may present signs of acute abdomen and shock indicating bowel infarction [2–4, 6].

The initial diagnostic examination of choice is ultrasound of the portal vein, which can show partial or complete filling of the vein with a solid isoechoic or hypoechoic material. Ultrasound has a sensitivity and specificity range of 80–100%. CT and angiography provide additional information on clot extent and presence of intestinal infarct that helps in the diagnosis with a sensitivity close to 90% [1–3]. Radiological features of acute versus chronic PVT are high luminal density in the
thrombosed vessel, lack of porto-portal collaterals, normal spleen size, no myeloproliferative disease, signs of intestinal wall edema, mesenteric vein anomaly, and filling defect [3].

As previously mentioned, in patients without liver cirrhosis or malignancy, PVT may occur due to inherited or acquired prothrombotic states. Inherited prothrombotic states are caused by genetic mutations. These mutations include factor V Leiden, prothrombin G20210A, and deficiencies of proteins C, S, and antithrombin III. Acquired prothrombic states are caused by changes in substances that affect coagulation and may be secondary to immobilization, trauma, surgery, pregnancy, malignancy, and medications. Antiphospholipid antibody syndrome, hyperhomocysteinemia, and myeloproliferative diseases are additional examples [7]. Medications that have been associated with prothrombotic states include hormonal therapies such as oral contraceptive pills, hormone replacement therapy (estradiol, progesterone, and testosterone), and tamoxifen (a selective estrogen receptor modulator). Additionally, anabolic steroids, methotrexate, erythropoietin, corticosteroids, chemotherapeutic agents, and cyclooxygenase 2 inhibitors have also been associated with elevated hypercoagulability [1, 6, 7].

Testosterone has been used for many years to improve libido in postmenopausal women [8, 9]. Testosterone pellets work by releasing small amounts of testosterone in the subcutaneous tissue, presumably over the span of 3–6 months. Testosterone pellets are
still not well regulated; for instance, they are very dif-
ficult to remove once inserted and can lead to levels
higher than those found in premenopausal women, for
which women need to be closely monitored. In 2019,
a global consensus position statement for the use of
testosterone therapy in women diagnosed with hypo-
active sexual desire disorder and sexual arousal dis-
order did not recommend the use of any preparation
that results in supraphysiologic concentrations of tes-
tosterone, including pellets and injections [10]. How-
ever, no significant association with an increased risk
of venous thromboembolism has been seen with TRT.
The etiology behind this increased risk is still unknown,
although it is hypothesized to involve interactions
with undiagnosed procoagulable factors [11]. Another
possibility is testosterone’s known effect stimulating
erthropoietin secretion, which leads to increases in
hemoglobin, hematocrit, and possibly blood viscos-
ity. Nevertheless, these events represent a low risk of
inducing prothrombotic events by themselves unless
other risks such as chronic hypoxemia as seen in sleep
apnea, obesity hypoventilation, COPD, or smoking
coeexist [1, 12, 13]. A study conducted to investigate an
association between secondary polycythemia and the
risk of venous thromboembolism did not show clearly
that polycythemia secondary to COPD or other chronic
hypoxic conditions independently increases the risk of
thrombosis. However, the authors did note that patients
in the study with venous thromboembolism had signifi-
cantly higher BMI [12]. The dangers of venous throm-
boembolism by testosterone most probably depends
on the patient’s age, weight, hormonal status, and addi-
tional prothrombotic risk factors.

Management of PVT involves investigating probable
causes such as malignancy, liver cirrhosis, hereditary
and acquired prothrombotic states, and treatment con-
sisting of anticoagulation with low-molecular-weight
heparin, heparin, and Coumadin, or new oral antico-
agulants. Additional treatment options are intravenous
thrombolysis and thrombectomy [2–5]. Intravenous
thrombolysis and thrombectomy can be considered if symptoms began within 12–18 hours before presentation. The treatment goal is to restore the portal vein circulation, thus preventing intestinal infarction and the eventual development of portal hypertension. Prompt treatment is essential because the thrombosed vessel rarely recanalizes spontaneously. Studies have shown favorable outcomes with early anticoagulation showing recanalization in the first week of diagnosis. Recanalization rates were complete in 38.3% and partial in 14% of patients [2–5]. The extent of the thrombosis, the presence of ascites, and prothrombotic states are negative predictors for recanalization. After treatment has begun, close monitoring for symptomatology indicating intestinal infarction is important. Improvement in abdominal pain and the presence of bowel sounds are indicators of good response to treatment. Persistent abdominal pain and bloating despite anticoagulation can indicate intestinal infarction, and prompt surgical exploration may be necessary to reduce mortality [2–6].

Studies have recommended anticoagulation for no less than 6 months in patients diagnosed with PVT with no specific cause identified. Long-term or lifelong anticoagulation may be recommended if the patient persists in a hypercoagulable state, has recurrent episodes, or has a personal or family history of deep venous thrombosis [2–5].

The prognosis for PVT is favorable if diagnosed early, and anticoagulation started promptly. The 5-year survival rate in this scenario is more than 85% [3, 11]. Complications are mostly related to either other predisposing conditions, late presentation, or surgery. The most feared complication is intestinal infarction because of its high morbidity and mortality of 20–50% [2, 3, 6].

In the present case, our patient started improving after day 2 of anticoagulation. Repeated abdominal CT revealed signs of portal vein recanalization (Fig. 2). The patient was later discharged on rivaroxaban for 6 months, was closely monitored with no recurrence and no evidence of portal hypertension, and was advised against any further TRT in the future.

**Conclusion**
We can conclude that, before starting TRT in postmenopausal women with hypoactive sexual desire, it is essential to do a thorough risk assessment for medical conditions as well as inherited and acquired prothrombotic states that may further increase the risk of venous thromboembolism. In addition, testosterone treatment should be tailored to the female physiologic testosterone levels and include close monitoring of pre- and post-treatment testosterone levels.

Hopefully, the benefits and risks of TRT for postmenopausal females will continue to be investigated. Finally, PVT is a rare but serious life-threatening event if not recognized and treated early. Therefore, the suspicion index should be high for clinicians evaluating severe abdominal pain in women on TRT.

**Abbreviations**
PVT: Portal vein thrombosis; HRT: Hormone replacement therapy; CT: Computed tomography, TRT: Testosterone replacement therapy; BMI: Body mass index, COPD: Chronic obstructive pulmonary disease.

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**Authors’ contributions**
MZC and LTOF collected patient data and prepared, evaluated, and edited the case report manuscript with literature review. MZC and LTOF revised and edited the abdominopelvic computed tomography scan images. MZC, LTOF, and ES were all involved in the patient treatment. ES participated in patient management and decision-making as a consultant from the specialty of Hematology/Oncology, and helped to review, edit, and add specific clinical insight to the manuscript. All authors read and approved the final version of the manuscript.

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**Competing interests**
The authors declare that they have no competing interests.

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