Duodenal intramural hematoma due to early postoperative anticoagulant treatment after a renal transplant: A case report

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ABSTRACT

A spontaneous intramural duodenal hematoma is a rare complication in patients receiving anticoagulation therapy. Presently described is a case of intramural duodenal hematoma in a patient with a cadaveric renal transplant who was under oral anticoagulant treatment due to paroxysmal atrial fibrillation. The patient was admitted with intense abdominal pain, nausea, vomiting, and a total obstruction of duodenum. After a diagnosis of intramural hematoma, a good prognosis was achieved with conservative care.

Keywords: Anticoagulant; intramural duodenal hematoma, renal transplantation.

INTRODUCTION

The intramural duodenal hematoma was first reported as “a false aneurysm” of the intestinal wall by McLauchlan during an autopsy in 1838.[1] An intramural duodenal hematoma usually occurs secondary to blunt abdominal trauma.[2,3] There are cases in the literature where an intramural duodenal hematoma occurs spontaneously and consequently the patient presents with abdominal pain due to partial gut obstruction. Spontaneous intramural duodenal hematomas are commonly associated with coagulopathy, anticoagulating drugs or sometimes with endoscopic procedures.[4–7] Warfarin is one of the usual suspects in these cases with a frequency of one case per 2500 anticoaugulated patients annually.[8]

CASE REPORT

Forty eight years old female patient had a cadaveric renal transplantation for end-stage renal disease secondary to hypertension and vesicoureteral reflux. She suffered early post-operative tachycardia diagnosed as paroxysmal atrial fibrillation which was managed with warfarin therapy. At the time of her discharge renal functions were normal and the International Normalised Ratio (INR) level was 2.1. Approximately 45 days after the transplant she was admitted to the emergency department with a two days history of epigastric pain, nausea and recurrent vomiting with no prior history of trauma. Abdominal examination was non-specific except for a relatively mild epigastric tenderness. Laboratory workup showed an increase in white cell count (14000 cells/mm³), but both liver and kidney functions were normal (creatine: 0.7 mg/dL). INR was 6.2 at the time and she was given fresh frozen plasma (FFP)/vitamin K combination to reduce INR levels. Plain graphs of the abdomen suggested intestinal obstruction. A nasogastric (NG) tube was inserted and 1500 cc bile colored intestinal fluid aspirated.

Upper gastrointestinal (UGI) water soluble contrast series showed minimal contrast passage to distal part and a collection of the barium in the 3rd portion of the duodenum (Fig. 1a, b). A subsequent UGI endoscopy confirmed dilatation of the 3rd portion of the duodenum and a total obstruction with an accumulation of drug tablets proximal to this level (Fig. 1c, d). CT scan of the abdomen/pelvis region with oral contrast demonstrated a non-enhanced, mildly hyperdense duo-
denal mass with a dilated intestinal segment proximal to this mass (Fig. 2a-d). The exact size of the mass cannot be measured due to the lack of IV contrast. Spontaneous intramural hematoma of the 4th portion of duodenum was suspected to be the cause and prolonged coagulopathy was thought to be the etiology of this situation.

After the initial diagnosis, decompression with a NG tube was obtained and a conservative treatment modality was followed through. In the following period, drainage from the NG tube decreased steadily. After removal of the NG tube, the patient was given standard oral feeding regimen which was tolerated without any problems.

In the end, no operative intervention was needed and obstructive symptoms resolved under conservative therapy without any complications. The patient was discharged with low molecular weight heparin treatment which was continued for six months. After an uneventful six months, the treatment ended and no additional intervention was needed in this time period.

**DISCUSSION**

Spontaneous intramural duodenal hematoma is commonly associated with coagulation abnormalities resulting secondary to usage of anticoagulant drugs. Oral anticoagulants such as warfarin are widely used and the most common complication is bleeding.[9] According to recent studies spontaneous intramural hematomas most commonly involve jejunum followed by ileum and duodenum; which is different than the traumatic intramural hematomas which mostly involves duodenum.[10]

A spontaneous intramural duodenal hematoma usually presents with intense abdominal pain followed by nausea and vomiting. Patients can be symptomatic for several days prior to presentation or diagnosis.[11–14] Several radiographic features have been described. These include a thickened intestinal wall on ultrasonography or circumferential wall thickening, intramural hyperdensity, luminal narrowing and intestinal obstruction on CT.[15,16] The problem is; these kind of radiologic abnormalities are not specific for intestinal tract hematomas and can resemble other pathologies.[16] So the clinician should have a low threshold in suspecting of a possible intramural hematoma diagnosis.

Apart from conventional USG and CT, UGI endoscopy is also a powerful tool in these cases. With the help of endoscopy, it is possible to both visualize and cure small-bowel hematomas. [17,18] Young Lee et al.[19] presented a case of an intramural
The conservative management consists of stopping oral anticoagulants immediately and starting a simultaneous infusion of FFP/vitamin K for reversal of anticoagulant effects. If the patient presents with obstructive symptoms; a NG tube should be administered swiftly and the drainage should be monitored closely.

Most cases have good prognosis with conservative care and the hematoma can start to regress as early as the first week. For management of spontaneous intestinal hematoma, there is no a global consensus on one specific approach. It usually responds well to conservative management within 10–15 days. [12,13] The conservative management consists of stopping oral anticoagulants and it is a manageable complication with conservative means.

A spontaneous intramural intestinal hematoma especially a duodenal one is a rare complication stemming from anticoagulating treatment modalities. Even though it is sometimes hard to diagnose properly, once the diagnosis is made it has a mild prognosis and long-term outcomes are good in the absence of any other concurrent conditions.

Conflict of interest: None declared.

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Böbrek nakli sonrası erken dönemde antikoagülan kullanılmakta bağlı gelişen duodenum intramural hematoma: Olgu sunumu

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Spontan duodenal hematoma, antikoagülasyon tedavi uyguladığı hastalarla nadir rastlanan bir komplikasyondur. Bu olgu örneğinde, kadavra duodenal hematoma, antikoagülasyon tedavi yaklaşım almakta olan hastada karşılaşılan spontan duodenal hematoma olgusunu sunulmaktadır. Kanın artsız, bulanti-kusma şikayetleriyle başvuran ve tam duodenal obstrüksiyon saptanan hastada spontan duodenal hematomun tanısal konulduktan sonra konservatif tedavi ile iyı bir prognoz sağlanmıştır.

Anahtar sözcükler: Antikoagülan; böbrek nakli; intramural duodenal hematoma.