Case report of unusual cause of lower GI bleeding

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Abstract
Lower GI bleeding is a challenging emergency for the gastroenterologist and the surgeon especially when it is massive and obscure in origin. We present a case of a 41 years old male who presented to our hospital with lower GI bleeding which was massive and of unknown origin. It necessitated taking the patient for surgical intervention. The patient was found to have a bleeding gastrointestinal stromal tumor (GIST) in the small bowel. Small bowels GISTs are rare and unusual to present with lower GI bleeding.

Case summary
41 years old gentleman known to have neurofibromatosis type 1 who, in the past, underwent surgery of frontal brain lobe xanthoastrocytoma presented to the local hospital with massive rectal bleeding where he was resuscitated and referred to our center for further investigations and management. The patient was haemodynamically reasonably stable when he was received in our centre with a hemoglobin of 9 g/dL. He continued to have rectal bleeding and an urgent upper gastrointestinal endoscopy and colonoscopy was arranged which did not identify the source of bleeding. Abdomen CT angiogram and then a mesenteric CT angiogram were performed but neither showed active contrast extravasation which could explain the source of bleeding. The CT showed extensive cutaneous and subcutaneous neurofibromatosis and left iliacus neurofibromatosis. The patient was kept on a high dependency unit where he received further 2 units of blood and his haemoglobin was stable around 9 g/dl. However, he became restless and drowsy and so he was intubated that stage and shifted to intensive care where he received further 2 units of blood and his haemoglobin did rise marginally from 8 g/dL to 8.2/dL, He continued to be off inotropes but tachycardic >110. So, the patient continued to pass P.R bleeding and was decided to be taken for surgical control. At laparotomy, there was a fairly vascular small bowel tumor 50 cm from ileocecal junction with no intraluminal blood distal to it. Another smaller one tumor was found just proximal to the first one (Figure 1). The first tumor was segmentally resected and continuity of the bowel was restored with side to side hand sewn anastomosis whilst the second tumor was locally excised. The patient remained stable in the postoperative period and made a swift recovery so he was discharged home after 5 days and seen in the clinic and found to have no complaints. The Histopathology of the resected tumors came back as gastrointestinal stromal tumor (GIST) low malignant potential. With good margins > 5 cm free of the disease. Then the Patient was referred to medical oncologist for further input. He was seen after 2 months from the surgery in the medical oncology with a CT of chest, abdomen and pelvis which was normal and started on imatinib and he still in Follow-up with them.

Discussion
There is a cohort of causes of lower GI bleeding which had been described in the medical literature (Table 1). These causes may vary with the age and other associated conditions. Our patient was a known case of neurofibromatosis type 1 (NF-1). This condition in particular may have some rare gastrointestinal manifestations.

NF-1 is inherited as autosomal dominant with complete penetrance and variable expression [1]. The frequency of intra-abdominal (gastrointestinal or retroperitoneal) manifestations of NF-1 varied greatly in previous studies ranging from 5%-25% [1].

GIST is a rare neoplasm accounting for about 1%-3% of all malignant neoplasms of the gastrointestinal tract. These tumors bear certain histopathological similarities to a specific cell type inherent in the GI tract, known as interstitial cells of Cajal (ICC) [2]. The majority of GIST cases (60% to 70%) arise from stomach. GIST tumours have been classified to myogenic, neurogenic and non-differentiated tumours [3].

Gastrointestinal manifestations of NF-1 usually arise during midlife or later; generally later than the appearance of the cutaneous manifestations of the disease. In a study from a large specialized centre,
intra-abdominal (gastrointestinal or retroperitoneal) manifestations necessitating surgical intervention occurred in 2.5% of patients, at a median age of 40 years, followed up regularly [3,4]. NF1 patients may be predisposed to developing small intestinal GISTs, which may appear as multiple GISTs [5].

The GIST tumour in our case was a rare manifestation of uncommon condition of neurofibromatosis as a cause of massive lower GI bleeding. It had been reported in the literature that such condition is rare cause of obscure lower GI bleeding. The cases reported were mainly jejunal GISTs which mainly presented as obscure lower GI bleeding long lasting for weeks [6]. Our patient was different in the sense that he presented acutely with significant lower GI bleeding that required a surgical intervention and his GISTs were in the ileum.

There are some reports that GISTs have been increasingly documented as an abdominal manifestation of the disease. GISTs were detected in 25% of NF-1 patients at autopsy [7,8]. Thus, they represent the most common gastrointestinal manifestation of NF-1 reported in literature.

However taken by site, NF-1 associated GISTs represented 6%, 4% and 0.06% of duodenal jejuno-ileal and gastric GISTs at the AFIP respectively. The mean age at GIST diagnosis was 49 years compared to a mean age of 56 years for small bowel GISTs in genera l. Most NF-1 associated GISTs present as small asymptomatic lesions with low mitotic activity and they generally follow a benign clinical course [8,9].

There has been a growing number of literature reporting an association between NF1 and GIST [10]. The malignancy is commonly determined by the tumor size and the mitotic ratio. Tumours larger than 5 cm in diameter or having a mitotic count greater than 5 per high-power field are classified as malignant [11-13]. In our case, the tumour was 3 cm in maximum diameter and with a low potential of malignancy as the mitotic figure was 5/50. Surgical resection is the treatment of choice. Lesions >2 cm should be resected. Lesions <2 cm can be monitored with serial imaging. Medical management involves treatment with Imatinib, which interferes with tumour proliferation by disrupting signalling via the C-kit oncogene [14].

Conclusion

Neurofibromatosis type 1 is an uncommon condition which may be present with lower GI bleeding and it should alert the surgeon for possibility of associated GIST tumor as a rare manifestation of such condition. There had been literature reports which described such lesions as a cause of lower GI bleeding [10-12], one of the rarest tumors is GIST. To our knowledge in this country (Oman) our case is the first case of such manifestation of NF and GIST of small bowel presented with massive lower GI bleeding.

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