A RARE FORM OF PRESENTATION OF A SMALL INTESTINE GIST IN THE EMERGENCY DEPARTMENT

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Abstract

Gastro-intestinal stromal tumors (GIST) are a form of mesenchymal tumors most commonly located in the gastro-intestinal tract, as the name suggests. The symptoms that might occur can be abdominal pain or discomfort, early satiety, bloating, jaundice, fatigue, hematemesis or melena. Most of GISTs (>95%) are positive for the KIT protein. Surgery remains the only curative treatment for GIST amendable for resection. High risk tumors can receive adjuvant or neoadjuvant treatment with Imatinib Mesylate, a tyrosin-kinase inhibitor. We present the case of a 59-years old patient that presented with rectal bleeding and abdominal diffuse pain. Blood tests shows leukocytosis and moderate anemia. CT scan describes an expansive, polycyclic process projected at the limit between the mesogastric region and the left flank, in close contact with the small intestine without any clear demarcation line between them and at the superior pole of the tumor, located in an enteral loop, a band of hyperdensity with net contour. Intraoperative, an extraluminal jejunal tumor was found, located at 15 cm from the duodeno-jejunal flexure, with intraluminal active bleeding, which lead us to suspect a GIST based on the macroscopic aspect. We performed segmental enterectomy of 10 cm of the small intestine containing the tumor with entero-enteral anastomosis end to end. The histopathological and immunohistochemistry examinations correlated with the macroscopic aspect have confirmed the diagnosis of GIST.

Keywords: GIST, gastro-intestinal stromal tumor, complicated GIST, hemorrhage

Introduction

Gastro-intestinal stromal tumor (GIST) is a mesenchymal tumor that was relatively recent discovered, usually being confused in the past with leiomyoma, leiomyosarcoma or leiomyoblastoma. From a morphological point of view GISTs could be epithelioid, spindle-cell or pleomorphic tumors. With the help of the evolution of the modern day immunohistochemistry and electronic microscopy, we can now say with certainty
that GISTs have a completely different kind of physiopathology: the activating mutation of the c-KIT tyrosine-kinase [1-3].

The main presentation is in the stomach (up to 70%) and small intestine (20-30%), but they can also occur in the esophagus, colon or rectum. Other locations like the omentum, mesentery or retroperitoneum are extremely rare [4,5].

GISTs has an extremely different potential of malignancy that could vary from small, benign tumors to very aggressive sarcomas.

The symptoms could vary depending on the location, the size of the tumor, so the patients could encounter abdominal pain or discomfort, early satiety, bloating, jaundice, fatigue, palpitation, hematemesis, melena [6]. The patients may also present with an abdominal mass and no other symptoms or, as other studies suggest, 10 to 25% of the patients present with metastases [7]. As some studies had shown, massive hemorrhage is rarely the first or main symptom that a patient can experience [8,9].

Classical therapeutic management is the surgical en-bloc removal of the tumor or, in case of metastatic disease, Imatinib Mesylate treatment prior to surgery. Depending on the location and size the removal of the GIST may require gastrectomy, pancreaticoduodenectomy or an abdominoperineal resection [3].

**Case presentation**

A 59-year old patient came to the Emergency Department with recurrent episodes of rectal bleeding and abdominal diffuse pain, symptomatology which started 7-day prior his admission and escalated quickly in the last 12 hours. The patient was known with cardiac failure NYHA II, high blood pressure, acute hemorrhagic gastritis, esophagitis caused by gastro-esophageal reflux disease, hepatic steatosis and cystic tumor in the liver’s second segment (confirmed by CT scan).

Clinical exam reveals pale complexion, diaphoresis and diffuse abdominal pain with signs of generalized abdominal guarding. Rectal examination indicates large quantity of melena.

Initial blood tests revealed leukocytosis (31.23 m/mm³) without any other sign of infection and moderate anemia (Hb = 8,1 mg/dL).

An abdominal and pelvic native and contrast CT was mandatory. In our patient’s case, this investigation showed an expansive, polycyclic process (67/49/81 mm) projected at the limit between the mesogastric region and the left flank, non-homogenous post contrast, with hypodense areas, in close contact with the small intestine without any clear demarcation line between them (Figure 1). At the superior pole of the tumor, located in an enteral ansae, a band of hyperdensity (39/9/3 mm) with net contour- most likely a foreign object (Figure 3). Furthermore, the CT scan reveals an adenopathic mass (26/10/46 mm) situated anterior to the inferior vena cava, adjacent to the caudate lobe and multiple adenopathies located in close contact to the small gastric curvature.

The patient was scheduled for emergency surgery under general anesthesia. Intraoperative, an extraluminal jejunal tumor with a diameter of 10 cm was found, located at 15 cm from the duodeno-jejunal flexure, with rich vascularisation, invasive in the serosa, with intraluminal active bleeding (Figure 4,5) which lead us to suspect a gastro-intestinal stromal tumor based on the macroscopic aspect. No other organs were affected. We performed segmental enterectomy of approximately 10 cm of the small intestine containing the tumor with entero-enteral anastomosis end to end (Figures 6,7).

The resected segment was sent for histopathological examination. The result describes the 7 cm tumor as a gastro-intestinal stromal tumor with fusiform pattern with a
mitotic index of < 5 mitosis / 5 mm², completely excised.

The immunohistochemistry examination shows positive results for CD117, DOG1, CD34, ACT and ki67. The histopathological and immunohistochemistry examinations correlated with the macroscopic aspect have confirmed the diagnosis of GIST, prognosis group 3a, medium risk of tumor progression – 24% AFIP.

The patient’s evolution was favorable and the clinical symptoms disappeared, remaining stable in the postoperative period and was discharged home in stable condition.
Gastro-intestinal stromal tumors are known for their unspecified clinical presentation. They can be easily misdiagnosed as leiomyoma, leiomyosarcoma, neuroendocrine tumors, inflammatory polyps and many other benign or malignant lesions [16]. They can also be confused with abscesses as other studies have shown [17].

In our particular case the diagnostic suspicion dictated by the CT scan and blood tests was foreign object inserted in the intestinal wall causing an abscess. The characteristic features of gastro-intestinal stromal tumors described on the CT images may frequently lead to diagnostic errors, as other studies have shown [18]. In this situation the correct approach for diagnostic and treatment is exploratory laparotomy and resection of the tumor, as the Report of GIST Consensus Conference shows [19].

CT is very useful in the diagnosis of the tumor, but the determination of the malignancy in GIST’s is difficult and cannot be resolved with use of imaging and histopathological features alone. The results of immunohistochemistry positive for CD117, DOG1, CD 34 are the correct choice in the diagnosis of gastro-intestinal stromal tumor [20].

The treatment for a severe bleeding GIST is emergency surgery, but the prognosis after resection is influenced by completeness of the excision (R0) and the malignant potential of such tumor. Adjuvant treatment may be needed for patients with incomplete resection or high grade GIST (Imatinib Mesylate). Complete surgical resection is considered sufficient for low grade GIST [21].

Conclusion

Even though gastro-intestinal stromal tumors are rare, they should always be considered in the differential diagnoses of hemorrhagic gastro-intestinal masses.
A rare form of presentation of a small intestine GIST in the emergency department

References


