Gastrointestinal Stromal Tumor in a Patient with Neurofibromatosis Type I: Diagnosis after Intestinal Occlusion Secondary to Adhesions after an Episode of Pancreatitis

doi: https://doi.org/10.32635/2176-9745.RBC.2023v69n2.3720

Tumor Estromal Gastrointestinal em Paciente com Neurofibromatose Tipo I: Diagnóstico após Oclusão Intestinal Secundária a Aderências após Episódio de Pancreatite

Tumor del Estroma Gastrointestinal en Paciente con Neurofibromatosis Tipo I: Diagnóstico tras Oclusión Intestinal Secundaria a Aderencias tras Episodio de Pancreatitis

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ABSTRACT

Introduction: Gastrointestinal stromal tumors (GIST), although relatively rare, account for 80% of mesenchymal tumors of the digestive tract. They manifest in any part of the alimentary tract and are derived from Cajal cells. They may occur sporadically or be associated with familial syndromes such as neurofibromatosis type I. The clinical picture is variable, and they are often diagnosed incidentally. The diagnosis requires imaging tests associated with histopathological and immunohistochemical analysis. The best strategy for treatment is surgical resection and cases should be analyzed individually to verify additional advantages with the association of systemic therapy. This study aims to present an unusual case of GIST associated with neurofibromatosis type I in a patient with incidental diagnosis after semientestinal occlusion secondary to an episode of pancreatitis, in addition to performing a literature review on the subject. Case report: A 49-year-old woman with a history of severe pancreatitis presented with intestinal obstruction approximately 8 months after this episode. Abdominal computed tomography revealed a heterogeneous formation in the mesogastrian region, measuring 6.6 x 5.1 x 5.3 cm. She underwent surgical resection and histopathological and immunohistochemical studies confirmed the diagnosis of GIST. Six months after diagnosis, the patient is in good general condition and is on systemic therapy. Conclusion: GIST are rare tumors, but their diagnosis should come to mind in patients with neurofibromatosis type I with abdominal masses.

Key words: gastrointestinal stromal tumors; neurofibromatosis 1; gastrointestinal neoplasms.

RESUMO

Introdução: Os tumores do estroma gastrointestinal (GIST), embora relativamente raros, correspondem a 80% dos tumores mesenquimais do trato digestivo. Manifestam-se em qualquer parte do trato alimentar e são derivados das células de Cajal. Pode ocorrer de forma esporádica ou associado a síndromes familiares como a neurofibromatose tipo I. O quadro clínico é variável, sendo frequentemente diagnosticados de forma incidental. O diagnóstico requer realização de exames de imagem associados à análise histopatológica e imunohistoquímica. A melhor estratégia para o tratamento é a reseção cirúrgica e os casos devem ser analisados individualmente para verificar vantagens adicionais com a associação da terapia sistémica. O objetivo deste trabalho é apresentar um caso inusual de GIST associado à neurofibromatose tipo I em uma paciente com diagnóstico incidental após semioclusão intestinal secundárias a episódio de pancréatite, além de realizar revisão de literatura sobre o assunto. Relato do caso: Mulher, 49 anos de idade, com passado de pancreatite grave, apresentou quadro de oclusão intestinal cerca de oito meses após esse episódio. A tomografia computorizada de abdome revelou formação heterogênea em região mesogastrian, medindo 6,6 x 5,1 x 5,3 cm. Foi submetida à reseção cirúrgica e os estudos histopatológico e imuno-histoquímico corroboraram o diagnóstico de GIST. Seis meses após o diagnóstico, a paciente encontrava-se em bom estado geral e em uso de terapia sistémica. Conclusão: Os GIST são tumores raros, porém seu diagnóstico deve ser lembrado em pacientes com neurofibromatose tipo 1 apresentando massas abdominais.

Palavras-chave: tumores do estroma gastrointestinal; neurofibromatose 1; neoplasias gastrointestinais.

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INTRODUCTION

Gastrointestinal stromal tumors (GIST) correspond to 1% of gastrointestinal neoplasms\(^1,2\). Although rare, they represent 80% of mesenchymal tumors of the digestive tract\(^3-7\) and 5% of all sarcomas. They can occur in any part of the alimentary tract, originating from the cells of Cajal, in the gastrointestinal myenteric plexus\(^2,4,6,8\). They are more common in the stomach (50-70%) and small intestine (20-30%), less frequently affecting the colon and rectum (2-5%) or other abdominal cavity organs such as pancreas, gallbladder, mesentery, omentum and peritoneum\(^3,4,7\).

Neurofibromatosis type I (NF1) has an autosomal dominant inheritance caused by mutations of the NF1 gene, located on chromosome 17q11.2.\(^8,9\) It is characterized by a wide range of manifestations and may be present in different parts of the body, including subcutaneous, bone, nervous and soft tissues\(^2,9,10\). Presumptive diagnosis is made through clinical criteria, and identification of the three main signs: neurofibromas, café au lait spots, and Lisch nodules, occurring in more than 90% of patients until puberty\(^10\).

Genetic disorders involved in NF1 is forty-five-fold more likely to develop a GIST\(^2\), in addition to other benign and malignant neoplasms, including neurofibroma, optic pathway glioma, malignant peripheral nerve sheath tumors, neuroblastoma, pheochromocytoma, and breast cancer\(^5\).

The clinicopathological features of GIST associated with NF1 are different from those of sporadic GIST. Patients are younger (mean age 49 years), lesions are multiple, occur primarily in the small intestine, are smaller in size, have low mitotic activity, and rarely exhibit mutations in the CD 117 (KIT) and Platelet-Derived Growth Factor Receptor Alpha (PDGFRA) genes. These patients lack specific mutations for GIST, therefore, they show a variable and usually incomplete response to treatment with reverse transcriptase inhibitors\(^6,11\).

Recently, specific treatments for rare subgroups of GIST, such as those associated with NF1, have appeared in the literature, contributing to the improvement both of diagnosis and management of this condition\(^12\). This study aims to present an unusual case of GIST associated with NF1 in a patient with an incidental diagnosis after adhesions secondary to an episode of pancreatitis, resulting in intestinal occlusion, in addition to literature review to contribute to the current literature.

The study complied with Resolution 466/2012\(^13\) of the National Health Council and was initiated after approval by the Institutional Review Board of “Universidade Federal de Sergipe”, CAAE (Submission for Ethical Review) number 63819322.9.0000.5546, and report number 5793112.

CASE REPORT

A 49-year-old woman presented to the University Hospital of “Universidade Federal de Sergipe”EBSERH, Aracaju, with abdominal pain associated with vomiting, constipation and weight loss of 12 kg. She had an acute necrohemorrhagic pancreatitis eight months before and abdominal CT scan demonstrated a solid lesion in the left epigastrium. She was referred to the surgical department for further investigation. The patient turned asymptomatic after clinical measures initiated at the hospital of origin. Neurofibromas and café au lait spots were observed on the entire body surface (Figure 1).

The abdomen was flat, soft, normal to percussion, and painless. A new CT scan was acquired and showed a heterogeneous formation in the left anterior region of the mesogastrium, in close relationship to the left rectus abdominis muscle and small bowel, with intermingled hypodense areas, measuring 6.6 x 5.1 x 5.3 cm (volume of 92.7 cm\(^3\)) (Figure 2).

Findings during exploratory laparotomy confirmed a mass adhered to the wall of jejunum and adhesions causing partial obstruction. The patient underwent partial intestinal resection and reconstruction with end-to-end anastomosis, and resection of a skin nodule. Macroscopy described a tumor measuring 7.0 x 5.1 x 4.6 cm adhered
GIST in a Patient with Neurofibromatosis Type I

Figure 2. Computed tomography showing heterogeneous formation in the left anterior abdomen

Figure 3. Immunohistochemical study characterizing positivity for CD 117 (c-KIT)

The patient evolved uneventfully and was discharged on the fifth postoperative day. After 14 months of follow-up, she remains in good general health, using Imatinib 400 mg/day.

DISCUSSION

GIST patients develop variable and nonspecific clinical presentation, according to size and location of the tumor, but are often asymptomatic and end up being diagnosed incidentally as in the present case. The most frequent symptoms are anemia, weight loss, gastrointestinal bleeding, abdominal pain, symptoms related to mass effect, and may also present intestinal obstruction, perforation, and peritonitis.

Cells of Cajal and GIST share common markers for CD 117 (also called the KIT gene) and a calcium-activated chloride channel called DOG. KIT is a transmembrane tyrosine kinase receptor, responsible for several cellular functions, including proliferation, adhesion, apoptosis, and cell differentiation. In GIST, mutation in the KIT gene is responsible for inappropriate activation of KIT, which causes oncogenic signaling in the cell without opposition to cell proliferation.

Other mutations can occur affecting succinate dehydrogenase (SDH), BRAF, KRAS, and NF1. Mutations in NF1 lead to formation of GIST in patients with NF1. While the two conditions may seem distinct, evidence suggests that NF1 gene may lead to increased risk of GIST. Some literature reports suggest that patients with NF1 should be monitored for GIST, especially if abdominal pain, vomiting, or gastrointestinal bleeding have been reported. Individuals having mutation in the NF1 have a probability of GIST of 7%. Conversely, NF1-associated GIST cases account for only 5% of all GIST. The molecular mechanism associating NF1 and GIST is still not completely clear. Such cases may be due to activation of the RAS/RAF/MAP kinase signaling pathway through loss of function of the NF1 tumor suppressor gene.

As for treatment, some authors claim that early detection and management of GIST in patients with NF1 can improve their outcomes.

GIST has been classified as to its risk of recurrence after resection, based on two prognostic factors: tumor size and mitotic count. The present case high-risk classification justifies adjuvant treatment with Imatinib.

Imaging techniques can be applied to help the diagnosis. On CT, these tumors are encapsulated and have heterogeneous contrast uptake due to the presence of areas of necrosis. It is also possible to identify areas of hemorrhage, intrallesional calcification, and metastatic disease. Metastases are not rare and the most affected sites include the liver and peritoneum. Magnetic Resonance Imaging can demonstrate pelvic, mesenteric and peritoneal extension.

Positron Emission Tomography improves diagnostic accuracy of metastases, and monitors clinical response to pharmacological treatment. It can detect lesions of up to 1 cm and epiploic metastases that go unnoticed on CT. It also provides information on metabolic activity and allows the quantification of the degree of malignancy, as greater the uptake of glucose by the tumor, greater is the metabolic activity and the aggressiveness.

On endoscopic examination, they appear as smooth submucosal tumors that protrude extrinsically into the visceral lumen and can sometimes ulcerate. Its location in the third and fourth layers (submucosa and muscularis propria) turns endoscopic biopsies inconclusive.

Endoscopic ultrasound identifies the layers of the digestive tract with great precision, allowing accurate diagnosis of submucosal lesions. However, it does not distinguish between GIST and another neoplasm of the intestinal wall, such as leiomyoma. Thus, fine needle aspiration is recommended for the diagnosis of the lesion.
Preoperative biopsy is not mandatory in suspected GIST, but it avoids the need for lymphadenectomy, which would be crucial for neuroendocrine tumors or intestinal adenocarcinoma. In addition, it is useful when the submucosal lesion is indeterminate or there is evidence of metastatic disease, and palliative treatment is considered.

Histopathological analysis may reveal the presence of fusiform (70%), epithelioid (20%) or mixed (10%) cells. Unlike other mesenchymal tumors, GIST expresses CD 117 antigen, demonstrated by immunohistochemical analysis.

The best treatment strategy is surgical resection. Routine lymph node resection is not performed, since metastases are rare. Lesions with suspected invasion of adjacent organs should be treated by radical monobloc resection. Meticulous surgical technique is required to prevent tumor/capsule rupture resulting in dissemination and worst prognosis.

Systemic therapy is indicated in case of metastases, non-resectable, or locally advanced tumors. Imatinib, the best-studied systemic agent, is an oral inhibitor of tyrosine kinase, c-KIT receptor and PDGFRα receptor. In general, the presence of KIT mutation is highly associated with response to this agent.

In the present case, the patient was symptomatic and presented some characteristics that have been associated with GIST in patients with NF1, such as younger age, and a localized isolated lesion in the small intestine, showing high mitotic activity, upon histological evaluation demonstrated specific mutations of sporadic GIST, contrary to what has been reported in the literature. Markers KI-67, DOG-1, CD-34 and CD-117 (C-KIT) were positive. Preoperative biopsy was not performed because the lesion was considered resectable, and suspicion of GIST was high.

CONCLUSION

GIST are rare tumors, with nonspecific symptoms, but their diagnosis should always be kept in mind when evaluating patients with NF1 and abdominal masses. The ideal treatment involves surgical resection with free margins and the use of Imatinib should be individualized.

CONTRIBUTIONS

Pamela Viana e Silva and Carlos Anselmo Lima contributed to the study design, in addition to data collection and critical analysis of the results. Pamela Viana e Silva, Thiago Menezes Costa, and Renanna Najara Veras Rodrigues contributed to the wording of the manuscript. All the authors revised and approved the final version for publication.

DECLARATION OF CONFLICT OF INTEREST

There is no conflict of interests to declare.

FUNDING SOURCES

None.

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