Laparoscopic antral resection with Billroth I reconstruction for a gastric glomus tumor

Hamzeh M. Halawani *, Mohammad Khalife, Bassem Safadi, Khaled Rida, Fouad Boulos, Farah Khalifeh

Department of General Surgery, American University of Beirut Medical Centre, Riad El Solh, Beirut 1107 2020, Lebanon

A R T I C L E  I N F O

Article history:
Received 24 June 2014
Received in revised form 11 September 2014
Accepted 8 October 2014
Available online 13 November 2014

Keywords:
Laparoscopy
Glomus Tumor
Antrum
Diagnosis

A B S T R A C T

INTRODUCTION: Gastric glomus tumors are fairly uncommon and mostly benign, with an estimated incidence of 1% of all GI soft tissue tumors. The most common GI site of involvement is the stomach, and in particular the antrum. Some cases have been discovered incidentally, but most are symptomatic presenting with GI bleeding, perforation or abdominal pain. Glomus tumors are submucosal tumors and hence mistaken for the more frequent gastrointestinal stromal tumors.

PRESENTATION OF CASE: A 33-year-old woman presented with intermittent dull upper abdominal pain for two days. Abdominal computed tomography (CT) was performed showing a hyperdense mass in the antrum. Endoscopy and endoscopic ultrasound revealed a submucosal antral mass along the greater curvature, suspicious for a gastrointestinal (GI) stromal tumor (GIST), a laparoscopic antrectomy with Billroth I reconstruction was done. Pathological examination revealed that the mass was a gastric glomus tumor.

DISCUSSION: The presented case report met all the usual standard criteria commonly used to identify glomus tumors, the uniqueness of the case lies in the occurrence of the glomus tumor in the stomach, first suspected as GIST, then confirmed as a gastric glomus tumor. The vast majority of glomus tumors of the GI tract have been described in the gastric antrum. They occur in adults of all ages with a significant female predominance (78%).

CONCLUSION: This case may aid in improving the recognition and diagnosis of this rare entity and in differentiating it from more common GISTs and gastric carcinoids. A built up knowledge between physicians is extremely necessary to avoid common confusion in taking the right medical approach.

© 2014 The Authors. Published by Elsevier Ltd. on behalf of Surgical Associates Ltd. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/3.0/).

1. Introduction

Glomus tumors are rare mesenchymal neoplasms composed of modified smooth muscle cells, closely resembling perivascular glomus bodies, with an occurrence of 1% among all soft tissue tumors. Though most commonly benign, they have been considered malignant in some rare cases. Glomus tumors are mostly found in distal extremities, but have been reported in the gastro-intestinal (GI) tract. The most common GI site of involvement is the stomach, and in particular the antrum. Some cases have been discovered incidentally, but most are symptomatic presenting with GI bleeding, perforation or abdominal pain. Glomus tumors are submucosal tumors and hence mistaken for the more frequent gastrointestinal stromal tumors (GISTs). We present a case of a symptomatic gastric glomus tumor that was treated with laparoscopic antrectomy and Billroth I reconstruction.

2. Presentation of case

A 33-year-old female presented to the emergency unit (EU) at the American University of Beirut Medical Center (AUBMC) complaining of dull intermittent upper abdominal pain that radiated to the right flank and groin over a two-day period. No symptoms of nausea, vomiting, change in bowl habits or bleeding per rectum were reported. Direct and rebound tenderness were noted in the right lower quadrant with a negative Giordano’s sign. Laboratory tests were unremarkable. A computed tomography (CT) scan of the abdomen and pelvis showed a 2.5 cm × 2 cm hyperdense gastric lesion (Fig. 1). Upper GI endoscopy and Endoscopic ultrasound (EUS) confirmed the presence of a submucosal antral...
mass along the greater curvature, suspicious for a GIST, hence a preoperative biopsy was not performed (Fig. 2). She was referred for surgical consultation and laparoscopic antrectomy with Billroth I reconstruction was performed because the mass was relatively large in size for wedge resection. Billroth I was preferred because tumors in certain parts of the stomach may require Billroth I or II to preclude any primary closure and to achieve gastrointestinal continuity (Fig. 3). Gross examination demonstrated a 2 × 1.6 × 1.6-cm submucosal and well-circumscribed mass (Fig. 4) and histology revealed a tumor consisting of a proliferation of round and extremely uniform cells arranged around vascular spaces of varying sizes. There was neither mitotic activity nor necrosis. No spindle cell component was identified. Immunohistochemistry confirmed that the mass stained strongly positive for smooth muscle actin (SMA), weakly positive for synaptophysin, and was negative for chromogranin and cytokeratin establishing the diagnosis of glomus tumor of the gastric antrum (Figs. 5, 6). The post-operative course of the patient was smooth. Feeding was started on day 3 post surgery and increased gradually. The patient tolerated the procedure very well with no complications throughout a follow-up period of 6 months.

3. Discussion

Glomus tumors of the GI tract are rare, representing around 1% of GI soft tissue tumors. They are defined as mesenchymal tumors composed of a population of modified smooth muscle cells with prominent vascular channels. The bulk of glomus tumors often originate in the neuromyoarterial glomus, an arteriovenous shunt that is supplied with nerve fibers performing a temperature-regulating function. Nearly most of the reported glomus tumors were located in the distal extremities, but they were exceptionally found in the GI tract. The vast majority of Glomus tumors of the GI tract have been described in the gastric antrum. They occur in adults of all ages with a significant female predominance (78%). Patients may present with ulcer-like symptoms such as epigastric pain, gastrointestinal bleeding or perforation in 31–35%, but some have been discovered incidentally intra-operatively or on routine endoscopy. Glomus tumors of the stomach are usually small with a median size of 2–3 cm.

Upper endoscopy generally reveals a well-defined subepithelial mass. Endoscopic ultrasound (EUS) findings may show a circum- scribed hypoechoic mass located in the third or fourth layer with a heterogeneous pattern. On CT imaging, glomus tumors display dense, homogeneous patterns. Imaging studies, including EUS and CT are therefore limited in the diagnosis of Glomus tumors due to non-specific and overlapping features with GIST.

The diagnosis of Glomus tumor is established on pathological examination. Three glomus tumor subtypes are generally described; solid glomus tumors, glomangiomas, and glomangio- myomas. This distinction is purely histologic depending on the relative prominence of specific components (prominent vessels in glomangiomas and prominent smooth muscle bundles in glomangiomyomas.

---

Fig. 1. CT scan of the abdomen and pelvis with IV contrast showing 2.5 cm × 2 cm hyperdense gastric lesion suggesting intramural tumor. (A) Axial cuts (B) coronal cuts.

Fig. 2. (A) Round submucosal lesion noted at the pylorus. (B) Endoscopic ultrasonography (EUS) shows 1.7 cm × 2.5 cm slightly hyperechoic round lesion arising from the muscularis propria.
The more important distinction is one between benign and malignant glomus tumors. This generally relies on the presence of certain features such as cellular atypia, mitotic activity, spindle cell areas, or frank sarcomatous change. Such findings may be focal and subtle hence the need for careful histopathologic examination of this rare neoplasm.\(^1\) Other diagnostic considerations such as carcinoid tumor and lymphoma should be easily resolved by the alert pathologist through careful morphologic examination and immunohistochemical stains.

Miettinen et al\(^5\) reported on 32 cases of Glomus tumors of the GI tract that were referred to the Armed Forces Institute of Pathology (AFIP) over an 18-year period. About half of those cases had a different diagnosis before final review (i.e.: GIST, carcinoid, lymphoma).

Histologically, the tumors typically had a solid pattern of sharply demarcated, round glomus cells with prominent, mildly dilated pericytoma-like vessels. Immunohistochemically, all tumors were positive for smooth muscle actin, vimentin and calponin, and nearly all had net-like pericellular laminin and collagen type IV positivity. All tumors were negative for desmin and S-100 protein. All tumors lacked KIT expression and the GIST-specific mutations in the c-kit gene. Glomus tumors never stain positive for chromogranin and are only weakly positive for synaptophysin. One patient from Miettinen's series died of diffuse metastatic disease. While this patient's tumor lacked significant mitotic activity, it did manifest atypical features such as spindle cell growth and vascular invasion.\(^12\) The literature contains a few such reports of malignant glomus tumor arising in different parts of the GI tract.\(^13,14\)
4. Conclusion

In conclusion, glomus tumors of the stomach should be considered in the differential diagnosis of gastric submucosal tumors. They cannot be distinguished from GIST on endoscopy, EUS or CT scan. Histological findings with immunohistochemical stains will allow a definitive diagnosis. Their small size and benign nature make them amenable to laparoscopic resection. Due to their rarity, there are no solid guidelines for follow-up. Yet, a careful step-by-step medical plan must be put ahead of time before approaching a patient with a submucosal gastric mass carrying a potential for malignant behavior. It is mandatory to carefully examine the mass histologically and immunohistochemically, to reach a definitive diagnosis. The value of such reports is to remind both surgeons and pathologists about these rare tumors of the GI tract, their potential confusion with other tumors of entirely different derivation and management algorithms, and the need to carefully assess them for the presence of atypical features that may predict malignant clinical behavior. Correctly diagnosing and classifying glomus tumors of the stomach and intestines is an essential part of providing patients with appropriate medical care and avoiding problematic and possibly negative outcomes.

Conflict of interest

The authors declare that they have no competing interests.

Funding

The American University of Beirut Medical Center sponsored the case report, but the efforts made for publication were made by the authors of the case.

Ethical approval

Written informed consent was obtained from the patient for publication of this Case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

All surgical and clinical protocols were approved by the institutional review board (IRB) at the AUBMC and signed by the ethics committee with consent for meeting the guidelines of Lebanon’s national healthcare system and in accordance with Helsinki Declaration.

Author’s contributions

MK and BS performed all surgical procedures presented in the case and contributed to the study design. KR collected data. HH drafted the manuscript, FK edited the manuscript, FB added the histological part and revised the manuscript. MK and BS revised and finally approved the manuscript for submission.

References

1. Park JP, Park SC, Park CK. A case of gastric glomus tumor. Korean J Gastroenterol 2008;52(5):310–4.
2. Huang CC, Yu FJ, Jan CM, Yang SF, Kuo YT, Hsieh JS, et al. Gastric glomus tumor: a case report and review of the literature. Kaohsiung J Med Sci 2010;26(6):321–6.
3. Matevosian E, Brucher BL, Nahrig J, Feussner H, Huser N. Glomus tumor of the stomach simulating a gastrointestinal stromal tumor: a case report and review of literature. Case Rep Gastroenterol 2008;2(1):1–5.
4. Zissis D, Ziui-Serbetzoglou A, Glava C, Grammatoglou X, Katsamagkou E, Nikolaidou ME, et al. Glomus tumor of the stomach: a case report. J BUON 2008;13(4):581–4.
5. Chou KC, Yang CW, Yen HH. Rare gastric glomus tumor causing upper gastrointestinal bleeding, with review of the endoscopic ultrasound features. Endoscopy 2010;42(Suppl. 4):ES8–9.
6. Miettinen M, Paal E, Lasota J, Sobin LH. Gastrointestinal glomus tumors: a clinicopathologic, immunohistochemical, and molecular genetic study of 32 cases. Am J Surg Pathol 2002;26(3):301–11.
7. Lee HW, Lee JJ, Yang DH, Lee BH. A clinicopathologic study of glomus tumor of the stomach. J Clin Gastroenterol 2006;40(8):717–20.
8. Fang HQ, Yang J, Zhang FF, Cui Y, Han AJ. Clinicopathological features of gastric glomus tumor. World J Gastroenterol 2010;16(36):4616–20.
9. Baek YH, Choi SR, Lee BE, Kim GH. Gastric glomus tumor: analysis of endosonographic characteristics and computed tomographic findings. Dig Endoscopy 2013;25(1):80–3.
10. Christopher DM, Fletcher KKH, Fredrik M. World Health Organization Classification of Tumours of the Soft and Bone Tissue; 2013.
11. Folpe AL, Fanburg-Smith JC, Miettinen M, Weiss SW. Atypical and malignant glomus tumors: analysis of 52 cases, with a proposal for the reclassification of glomus tumors. Am J Surg Pathol 2001;25(1):1–12.
12. Xu XD, Lu XH, Ye GX, Hu XR. Immunohistochemical analysis and biological behaviour of gastric glomus tumours: a case report and review of the literature. J Int Med Res 2010;38(4):1539–46.
13. Gassel HJ, Klein I, Timmermann W, Kenn W, Gassel AM, Thiede A. Presentation of an unusual benign liver tumor: primary hepatic glomangiomatosis. Scand J Gastroenterol 2002;37(10):1237–40.
14. Abu-Zaed A, Azzam A, Amin T, Mohammed S. Malignant glomus tumor (glo- mangiosarcoma): of intestinal ileum: a rare case report. Case Rep Pathol 2013;2013:305321.

Open Access

This article is published Open Access at sciedirect.com. It is distributed under the IJSSCR Supplemental terms and conditions, which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original authors and source are credited.