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**Primary gastroduodenal tuberculosis presenting as gastric outlet obstruction: A case report and review of literature**

Ali AM *et al*. Primary gastroduodenal tuberculosis

Abdihamid Mohamed Ali, Yahye Garad Mohamed, Abdirahman Ahmed Mohamud, Abdulkadir Nor Mohamed, Mohamed Rage Ahmed, Ismail Mohamud Abdullahi, Tuba Saydam

**Abdihamid Mohamed Ali, Abdirahman Ahmed Mohamud, Abdulkadir Nor Mohamed, Mohamed Rage Ahmed, Tuba Saydam,** Department of General Surgery, Mogadishu Somali Turkey Recep Tayyip Erdogan Training and Research Hospital, Mogadishu 2526, Somalia

**Yahye Garad Mohamed,** Department of Radiology, Mogadishu Somali Turkey, Training and Research Hospital, Mogadishu 2526, Somalia

**Ismail Mohamud Abdullahi,** Department of Pathology, Mogadishu Somali Turkey Recep Tayyip Erdogan Training and Research Hospital, Mogadishu 2526, Somalia

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**Corresponding author: Abdihamid Mohamed Ali, MBChB, MD, MMed, Academic Editor, Academic Research, Lecturer,** Department of General Surgery, Mogadishu Somali Turkey Recep Tayyip Erdogan Training and Research Hospital, Hodan Street, Mogadishu 2526, Somalia. abdihamidmohamed10@gmail.com

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**Abstract**

BACKGROUND

Mycobacterium tuberculosis (TB) is the causative agent of TB, a chronic granulomatous illness. This disease is prevalent in low-income countries, posing a significant global health challenge. Gastrointestinal TB is one of the three forms. The disease can mimic other intra-abdominal conditions, leading to delayed diagnosis owing to the absence of specific symptoms. While gastric outlet obstruction (GOO) remains a frequent complication, its incidence has declined with the advent of proton pump inhibitors and *Helicobacter pylori* eradication therapy. Gastroduodenal TB can cause upper gastrointestinal hemorrhage, obstruction, and malignancy-like tumors.

CASE SUMMARY

A 23-year-old male presented with recurrent epigastric pain, distension, nausea, vomiting, and weight loss, prompting a referral to a gastroenterologist clinic. Endoscopic examination revealed distorted gastric mucosa and signs of chronic inflammation. However, treatment was interrupted, possibly owing to vomiting or comorbidities such as human immunodeficiency virus infection or diabetes. Subsequent surgical intervention revealed a dilated stomach and diffuse thickening of the duodenal wall. Resection revealed gastric wall effacement with TB.

CONCLUSION

Primary gastric TB is rare, frequently leading to GOO. Given its rarity, suspicions should be promptly raised when encountering relevant symptoms, often requiring surgical intervention for diagnosis and treatment.

**Key Words:** Tuberculosis; Gastrointestinal tuberculosis; Gastric outlet obstruction; Gastroduodenal tuberculosis; Case report

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**Core Tip:** Tuberculosis (TB) is a disease that has been around since ancient times and is still one of the top 10 causes of death globally. With a high incidence in lower-income countries. It is a global health concern, with gastrointestinal TB The third most common extrapulmonary form of TB. Here we present Primary gastroduodenal TB presenting as gastric outlet obstruction (GOO): A case report and review of the literature. The diagnosis of abdominal TB is often delayed since there are no distinct clinical signs and symptoms, and the disease can mimic other intra-abdominal pathologies. The most common causes of GOO include peptic ulcer disease and gastric cancer. TB should be included ddx when these dx is excluded. Surgery is inevitable when GOO is present at time of diagnosis and proceed by antitubercular treatment.

**INTRODUCTION**

Tuberculosis (TB) is a chronic granulomatous disease caused by aerobic bacteria *Mycobacterium* TB[1,2].

Its global incidence is high in low-income nations, posing a significant health concern. TB has high morbidity and mortality rates despite its status as a treatable disease[3]. Abdominal TB ranks as the third most common extrapulmonary form[4].

Gastrointestinal TB (GI–TB) is one of the three forms of abdominal TB. The other types include visceral, peritoneal, and tuberculous lymphadenopathy[5].

*Mycobacterium* TB enters the gastrointestinal system *via* hematogenous spread, ingestion of contaminated sputum, or direct dissemination from infected adjacent lymph nodes and fallopian tubes[6,7].

Diagnosing abdominal TB is often hindered by the absence of distinct clinical signs and symptoms, leading to delayed identification because the disease can mimic other intra-abdominal pathologies[8,9].

Correcting the general status of patients with a confirmed diagnosis of GOO, administering intravenous fluids to correct an electrolyte imbalance, and performing gastric decompression by inserting a nasogastric tube are all essential to the initial management of these patients[10]. The most common causes of gastric outlet obstruction (GOO) are peptic ulcer disease (PUD) and gastric cancer. However, the frequency of GOO owing to PUD has declined since the development of proton pump inhibitors and Helicobacter eradication therapy.

Gastric bezoars, pancreatitis-related fluid collection, caustic ingestion, massive gastric polyps, Crohn's disease, and complications following gastric surgery contribute to GOO[11].

Gastroduodenal TB (GD-TB) presents with various manifestations, including upper gastrointestinal hemorrhage, obstruction, and gastric or periampullary tumors suggestive of malignancy[12].

Similar to our patient, most individuals with GD-TB exhibit signs of GOO, often attributed not only to intrinsic duodenal lesions but also to extrinsic compression by tuberculous lymph nodes[13].

Here, we present a case of GD-TB-associated GOO in a previously healthy 23-year-old patient who exhibited no signs of pulmonary TB.

**CASE PRESENTATION**

***Chief complaints***

A 23-year-old male patient came to the Department of Surgery with recurrent epigastric pain, vomiting, and weight loss that had persisted for a year despite the patient having no prior history of chronic illness.

***History of present illness***

The patient had a year-long history of recurrent epigastric pain, distension, low appetite, nausea, vomiting, and weight loss. He did not mention any fever, jaundice, or changes in bowel habits. Hemostasis and cough were absent, and there were no notable findings from the assessment of the other systems.

***History of past illness***

The patient had experienced similar symptoms during a prior stay in South Africa eight months earlier, during which an endoscopy revealed distortions and chronic inflammatory changes in the distal gastric mucosa without evidence of *Helicobacter pylori (H. pylori)* infection. Subsequently, anti-tubercular medication was planned, but the patient ceased treatment after one week owing to intractable vomiting.

***Personal and family history***

There was no family history of similar conditions or TB.

***Physical examination***

The patient presented with normal vital signs, absence of pallor, and no cervical, axillary, or inguinal lymphadenopathy.

Clear chest examination findings were noted. Upon abdominal examination, the abdomen was soft and non-tender with mild distension, and no masses, organomegaly, or ascites were noted.

***Laboratory examinations***

C-reactive protein 200 mg/L, stool *H. Pylori*-negative. No abnormalities were found in routine blood and urine analyses.

***Imaging examinations***

An abdominal contrast-enhanced computed tomography (CT) scan revealed gastric distension and diffuse wall thickening in the first and second parts of the duodenum. A thoracic CT scan showed no evidence of prior TB sequelae (Figure 1). Upper endoscopy was subsequently performed, revealing longitudinal hyperemic mucosal streaks in the gastric corpus and antrum, along with multiple glandular nodules and severely inflamed mucosa in the post-pyloric duodenum. Additionally, obstruction was observed at the junction of the first and second parts of the duodenum.

A gastric biopsy revealed chronic gastritis with severe inflammation, absence of metaplasia and dysplasia, and a positive result for *H. pylori*.

The patient was admitted for optimization of preoperative fitness in preparation for surgery.

**FINAL DIAGNOSIS**

Primary GD-TB.

**TREATMENT**

A decision was made to relieve the obstruction surgically. Intraoperatively, the following findings were noted: Dilation of the stomach and diffuse wall thickening of the duodenum, along with multiple enlarged mesenteric lymph nodes (located at the lesser curvature, transverse colon mesentery, small bowel mesentery, paraduodenal area, and interaortocaval region). Subsequently, antrectomy and Roux-en-Y reconstruction with gastrojejunostomy were performed, along with lymph node dissection (Figure 2).

The pathology report from the resected distal part of the antrum and lymph nodes revealed findings indicative of gastric TB, characterized by effacement of the gastric wall architecture and numerous caseating granulomatous inflammation. Specifically, the distal margins displayed positivity for granulomatous inflammation. Additionally, 12 Lymph nodes exhibited suppurative and non-suppurative granulomatous inflammation, further supporting the TB diagnosis. Periodic Acid-Schiff stain stains yielded negative results for fungal hyphae. These findings were consistent with GD-TB (Figure 3). Notably, cytology analysis for Acid Fast Bacteria staining and Gene Xpert testing were not pursued owing to the absence of peritoneal ascites detected during the operation.

**OUTCOME AND FOLLOW-UP**

The postoperative course was uneventful, and the patient was discharged from the hospital following successful tolerance of full enteral feeding. Plans were made for follow-up visits at the polyclinic and referral to a TB center to initiate anti-tubercular treatment. Currently, the patient is adhering to the anti-tubercular medication regimen well and has experienced no adverse effects.

**DISCUSSION**

GI–TB most frequently affects the ileocecal area, with the colon and jejunum following closely after. A total of 64% of GI–TB cases are caused by jejunal and ileocecal TB[14]. However, the esophagus, stomach, and duodenum are rarely affected. The duodenum and gastric are the primary sites of involvement for gastric TB.

Primary and isolated stomach TB is exceptionally rare, documented only in very few cases in the literature. A total of 0.4%–2% of all GI–TB cases are associated with primary gastric TB, while 2%–2.5% are associated with primary duodenal TB[15].

Several factors contribute to the low frequency of gastric TB, including the bactericidal properties of gastric acid, the presence of thick and intact gastric mucosa, and the absence of lymphoid structures in the gastric mucosa[14,16]. Treatment with H2 blockers increases the likelihood of involvement of the lesser curvature and pylorus in gastric TB, often presenting as ulcerating lesions[5]. Manifestations such as pyloric stenosis, miliary tubercles, and hypertrophic variations may also occur[14]. Gastric TB is frequently associated with TB lymphadenitis, as observed in cases where peripancreatic lymph nodes are involved, alongside visible ulcerative lesions in the prepyloric region.

The third part of the duodenum is commonly affected in cases of primary duodenal TB.

Both intrinsic and extrinsic factors can contribute to duodenal involvement[17]. Extrinsic involvement is most prevalent and is often related to lymphadenopathy in the duodenum's C-loop. Intrinsic variations may manifest as ulcerative, hypertrophic, or ulcer hypertrophic forms, potentially leading to fistula or stricture formation.

Various modalities contribute to GI–TB involvement, including ingestion of contaminated milk or food (primary TB), ingestion of contaminated sputum (secondary TB), hematogenous spread from a distant TB focus, or contiguous dissemination from infected neighboring foci *via* the lymphatic channels[18,19]. Given the absence of evidence of extra-abdominal TB in our case, we believe the infection to be primary gastric TB involving the peripancreatic lymph node.

GD-TB presents with nonspecific clinical characteristics commonly associated with weight loss, epigastric pain, and fever. In certain cases, GOO may be the presenting feature. Gastric TB may also mimic lymphoma or carcinoma, complicating the differential diagnosis. A study by Rao *et al*[12] conducted at a single center in India reported that among 23 patients with histologically confirmed GD-TB, vomiting (60.8%) and epigastric pain (56.5%) were the most prevalent presenting symptoms, with characteristics of GOO observed in 61% of cases (14 patients)[12]. While two patients had pyloric stenosis, the remaining twelve experienced obstruction owing to duodenal stricture. Additionally, out of the 23 patients studied, four had diabetes mellitus, and none were human immunodeficiency virus-positive. Similar to this study, our patient presented with vomiting, epigastric pain, and early satiety, suggestive of GOO.

The lack of specific clinical symptoms and diagnostic indicators often leads to underdiagnosis of GD-TB.

Histopathological examination of gastroduodenal biopsy specimens obtained *via* endoscopy remains the gold standard for diagnosing GD-TB. However, owing to the submucosal nature of granulomatous lesions, they are often challenging to identify, even in biopsy samples. A review revealed that granulomas were detected in only seven out of 27 patients who underwent endoscopic biopsy for duodenal TB[20].

Similarly, Rao *et al*[12] reported positive biopsies in their study in only 2 out of 20 patients.

Complications of GD-TB may include GOO, hemorrhage, and perforation, which can significantly increase morbidity[11].

As outlined in a review by Rao *et al*[12], in cases of TB-related GOO, truncal vagotomy and gastrojejunostomy (with or without feeding jejunostomy) were performed in twelve out of fourteen patients[12].

While abdominal TB can affect individuals of any age, there is a significant predominance among women, particularly those between the ages of 25 and 45[21]. In patients with GD-TB presenting with obstruction or mass, the yield of endoscopic biopsy is typically low[22,23].

Dyspeptic symptoms suggestive of gastric lesions often lead to suspicion of peptic ulcer, while weight loss may prompt consideration of gastric cancer as the primary diagnosis.

Up to 20% of patients undergoing examinations may exhibit evidence of pulmonary TB on chest X-ray[24], and duodenal bulb deformity may be detected during upper gastrointestinal endoscopy[25]. Because ulcerated lesions predominantly reside in the submucosa, conventional endoscopic biopsies often yield poor results and rarely uncover granulomas[21].

Although preoperative diagnosis of duodenal TB is exceedingly rare[17,26], Debi *et al*[14] have demonstrated that an endoscopic ultrasound is a valuable modality for characterizing lesions and obtaining samples for cytological confirmation[1]. In cases where histological diagnosis cannot be established by other means, intraoperative fine-needle aspiration cytology may be employed to obtain samples from the affected duodenal section[27].

Most lesions diagnosed with TB before surgery respond well to appropriate antitubercular treatment and may not require surgical intervention[28]. Antituberculosis medication therapy is the cornerstone of medical treatment for gastric TB. However, surgery becomes necessary for patients experiencing severe GOO owing to hypertrophic TB, with antituberculosis medication therapy following surgery[29].

In cases of GOO, gastrojejunostomy is preferred over pyloroplasty owing to the severe fibrosis around the pyloroduodenal junction, which may compromise the safety of pyloroplasty.

This study adheres to the surgical case report guidelines 2016 criteria[30].

**CONCLUSION**

Primary gastric TB is rare and often poses a diagnostic challenge, particularly when it presents as GOO. In regions where TB is prevalent, heightened suspicion is warranted. Surgery is often essential for diagnosis and treatment.

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**Footnotes**

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**Figure Legends**

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**Figure 1 Oral and intravenous contrast-enhanced thoraco-abdominal computed tomography scans.** A: It demonstrates gastric distension; B: It accompanied by multiple mesenteric lymphadenopathies; C: Duodenal wall thickening and nearly complete obstruction in the second part of the duodenum; D: Thoracic computed tomography findings indicated normalcy, with no active pulmonary tuberculosis or sequelae observed.

图片包含 游戏机, 食物, 披萨

描述已自动生成

**Figure 2** **Intraoperative finding.** A-C: An intense inflammatory mass causing duodenal obstruction alongside multiple pathologic lymph nodes observed in the mesenteric and paraduodenal regions; D: Resected distal gastric and dissected lymph nodes.

墙上挂着一幅画

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**Figure 3** **Histopathology.** A: Granulomatous inflammation and giant cells on the serosal surface; B: Lymph node parenchyma with effacement of multiple large tuberculoid necrotizing granuloma; C: Lymph node parenchyma with effacement of large tuberculoid necrotizing granuloma (thick arrows) epithelioid histiocytes (thin arrows) peripheral lymphocytes (triangles); D: Lymph node with tuberculoid necrotizing granuloma (thick arrows) and Langerhans giant cells (thin arrows) and peripheral lymphocytes (triangles).