Fungal Infections Masquerading as Gastrointestinal Tract Malignancies – A Series of Three Cases

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ABSTRACT
Fungal infections of the gastrointestinal tract (GIT) are not rare, but get missed due to their misleading clinical features. Here we report three GIT fungal infections at different locations with varied symptoms. We received surgical resection specimens with the clinical suspicion of malignancy for all. The first was a case of Aspergillosis, at an uncommon site, esophageal, in a 60-year diabetic male with a suspicion of carcinoma in view of presence of dysphagia and a stricture on CT. The second was a 60-year male, a case of stomach Zygomycosis with no definite immunosuppression, who presented with an intra-operative suspicion of malignancy, and showed fungal angio-invasion and ischemic perforation. Third was a 32-year immunocompetent male, a case of GI histoplasmosis with an uncommon presentation, showing obstructive symptoms due to an ileocecal mass with a clinical suspicion of TB/malignancy. Thus, a high index of suspicion for fungal a etiology in GIT is essential, because misdiagnosis can lead to a radical surgery. Timely treatment with appropriate antifungal therapy is crucial as high mortality is associated with certain fungi.

Keywords: Gastrointestinal Tract, Fungal Infections, Malignancy, Aspergillosis, Zygomycosis, Histoplasmosis

Case Report

Case 1: A 60-year-old man, a known case of diabetes mellitus presented with a history of dysphagia. His endoscopy and CT scan showed esophageal stricture with a possibility of malignancy. Repeated multiple biopsies performed...
showed only inflammatory exudate. However, in view of strong clinical suspicion of carcinoma, patient underwent esophagectomy with partial gastrectomy. We received an esophago-gastrectomy specimen, measuring 6.5 cm. The wall of esophagus was externally thickened and nodular, while the cut section showed a stricture with focal mucosal ulceration. No definite mass lesion was identified, either in esophagus or stomach. Routinely stained (H & E) sections (figure 1: a-c) showed ulcerated oesophageal mucosa with fibro-collagenous tissue in the wall, and multiple granulomas with giant cells and mixed inflammation, seen in muscularis propria and adventitia. Hence special stains were asked, of which Acid fast Bacilli (AFB) was negative and Gomori’s Methenamine Silver (GMS) stain showed acutely branched, focally septate fungal hyphae resembling Aspergillus. Multiple sections from esophagus and stomach showed no evidence of malignancy. Hence a diagnosis of “Esophageal granulomatous inflammation with stricture formation due to invasive Aspergillosis” was rendered. A culture was advised which subsequently came positive. Patient had indolent course postoperatively and was started on antifungal to which he responded well. A culture was advised which subsequently came positive. Patient had indolent course postoperatively and was started on antifungal to which he responded well.

Case 2: A 60-year-old man presented in emergency with acute abdominal pain with the features of perforative peritonitis. He was taken for emergency exploratory laparotomy, and a distal gastrectomy was done for gastric perforation with an intraoperative suspicion of malignancy. No significant past history could be elicited. Specimen received showed multiple large perforations, lined with black necrotic material, associated with an indurated and flattened mucosa with diffuse thickening of the stomach wall, resembling ? Linitis plastica (figure 2: a). H&E stained sections showed mucosal ulceration with inflammatory infiltrate associated with necrosis of mucosa and markedly congested blood vessels, few with fresh fibrin thrombi, suggesting ischemic changes (figure 2: b). However, also seen amidst inflammation on closure view were, scattered filamentous structures resembling fungal hyphae positive on GMS stain, which were broad, aseptate with wide angle branching, thus favouring Histoplasmosis. (figure 2: c). Few vessels also showed invasion by these hyphae with thrombus formation (figure 2: d). Thus a histological diagnosis of “Gastric ulcer with perforation due to fungal etiology, most likely mucor” was made, and an advice for a culture was given. Patient subsequently received antifungal treatment, but left against medical advice and was lost for any further follow up.

Case 3: A 32-year-old man presented with intestinal obstruction with complaints of distension of abdomen with colicky pain, multiple episodes of vomiting and constipation since last three days. He was a known alcoholic and smoker, negative for HIV, but with a past history of Anti-Koch’s Treatment (AKT) taken irregularly just for 3 months due to a clinical suspicion of pulmonary and abdominal tuberculosis. On physical examination, there were multiple bilateral posterior triangle Lymph Nodes, largest measuring 1.5 cm, firm, mobile, non-tender. On per abdomen examination, abdomen was distended, tense with palpable bowel loops. His X-ray showed multiple air fluid levels in abdomen, and mass in caecum on CT scan with a possibility of malignancy. Thus, a cervical lymph node FNAC was advised to rule out metastasis. Subsequently, quarter colectomy was performed, and specimen was received in segments, ileum measuring approximately 15 cms and ileo-caecal junction with ascending colon measuring 19cms, with no tubercles or strictures on external examination. On cutting open, a nodular, and soft to firm mass measuring 1 cm in diameter was seen in ileum, 2cm away from the resection margin, which showed greyish white, flashy, glistening cut surface and an adjacent perforated ulcer (figure 3: a, b). The caecum and appendix showed diffuse thickening and a similar cut surface on cutting open with necrotic material in Appendicular lumen (figure 3: c). Few enlarged mesenteric lymph nodes were also found with no caseation. Meanwhile, the FNAC report received showed, necrotizing granulomatous inflammation rich in histiocytes, with intra & extracellular yeast forms with single eccentric nucleus and surrounding “halo,” thus favouring Histoplasmosis. (figure 3: d). Histological sections from Ileum, caecum, appendix and mesenteric nodes also showed multiple granulomas with giant cells and histiocytes having foamy cytoplasm with numerous intracellular tiny dots like organisms, positive on GMS and PAS stain (figure 3: e-f), with narrow based budding. Thus a final diagnosis of “Disseminated Histoplasmosis with gastrointestinal perforation” was made.

Discussion

In this study, we came across a series of three cases of fungal infections of different species, involving different GI sites, but all presented with a clinical suspicion of a malignancy on CT/ intraoperatively/ on gross pathology. First case was angio-invasive aspergillosis involving esophagus, second was zygomycosis involving stomach, while third was a case of disseminated histoplasmosis involving ileum and caecum.

The importance of fungal infections of the GI tract has increased in parallel with the numbers of patients with organ transplants, AIDS, and other immunodeficiency states. Although fungal infections of the GI tract are often a part of a disseminated disease process, GI symptoms and signs may be the only presenting manifestations of disease. [7]
Fig. 1: a) Ulcerated esophageal mucosa (black arrow), dense inflammatory infiltrate, fibrocollagenous tissue and multiple granulomas (red arrow) in submucosa (H&E x 40). b) Muscularis propria showing granulomas with giant cells (H&E x 100) c) Granuloma with giant cells (H&E x 400) and slender fungal hyphae with acute angle branching on GMS stain (inset).

Fig. 2: a) Stomach showing Perforations, lined with black necrotic material, with diffuse thickening of the wall (inset). b) Ulcerated stomach mucosa, inflammatory granulation tissue, congested blood vessels; many with fibrin thrombi (arrow) (H&E x 40). c) Necro-inflammatory background with broad, pauci-septate fungal hyphae with wide angle branching (H&E x 400), positive on GMS. d) Angioinvasion, with destruction of arteriolar wall by fungal hyphae (arrow) (H&E x 400).
Fig. 3: a) External surface of ileum showing a nodular mass. b) Greyish white, flashy, glistening cut surface with an adjacent ulcer (arrow). c) Diffusely thickened caecum and appendix. d) Cervical lymph node FNAC showing granulomatous inflammation with extracellular (arrow) and intracellular (inset) yeast forms with eccentric nucleus and surrounding “halo” (MGG x 400). e) Ileum showing lymphohistiocytic inflammation with dot like organisms (arrow) (H&E x 40) and occasional granulomas (inset). f) Histiocytes showing intracellular tiny organisms (arrow) (H&E x400), positive on GMS stain (inset), with narrow based budding.
Invasive Aspergillosis (IA) typically involves the lungs, but may also infect the nasal sinuses, central nervous system, and rarely gastrointestinal (GI) system. It can be seen, both in immunocompromised and immunocompetent hosts. Mishra et al [2] and Cha et al [8] have both reported GIA in immunocompetent patient. Isolated GIA is very rare (approximately 5%) even in immunocompromised hosts. Eggiman et al, in a large study of GIA (n=1538), observed most cases in disseminated disease; isolated gut aspergillosis was very exceptional (0.8%). [9] In our study, the patient of GIA was though diabetic, but he did not show signs of any other organ involvement, thus possibly had isolated GI involvement.

The common presenting features of GIA are abdominal pain, fever, weight loss and perforation, and sometimes can even mimic malignancy similar to our case, which is similarly also reported by Mishra P et al in a case of gastric aspergillosis. [2] Most of the studies done on GIA, have shown lower GI tract as most common site involved (small bowel, colon). However, in certain common pathologic conditions like gastric ulcers and severe gastritis, the mucosal barrier can be penetrated by aspergillus spores and can lead to invasive gastric or upper GIA. [8] Kazan et al. in a review of 21 cases reported most cases involving lower GI tract, and only three from upper GI tract. [16] Further, the esophageal involvement by aspergillosis is uncommon as compared to candidiasis, and oedynophagia and dysphagia are usual presenting complaints [3] as seen in our patient too. The clinical manifestations at other sites are often nonspecific, such as abdominal pain, diarrhea, hemorrhage, and occasionally intestinal obstruction and perforation. Stricture however is not a common feature, [8] but was present in our case. The histological features of GIA are same as described at any other site, sometimes with a characteristic nodular infarction like lesion which was not present in our case. The inflammatory response ranges from minimal to marked, with a prominent neutrophilic infiltrate. However, a granulomatous response is not very common, but was quite pronounced in our case. [3]

Zygomycetes are filamentous fungi having low intrinsic pathogenicity. However, these fungi can produce fulminant infection in patients with underlying immune compromised conditions. Disease can be disseminated or localized to the rhinocerebral area, lungs, gastrointestinal tract or skin [11], and rarely may it affect healthy individuals; as is reported by Sunitha et al. [4] In our case too, there was no history suggesting any associated immunosuppression. Stomach is the most common site involved in GIZ, accompanied by a very high mortality rate. Other sites being colon followed by ileum. [4] The common clinical presentation of GIZ is massive gastric bleed, associated with non-specific symptoms like nausea, vomiting, abdominal pain or distension, depending upon the site of involvement. [12] Thus, sometimes the presentation can mimic inflammatory bowel disease as reported by Sunitha et al [4] and even malignancy as reported by Debata et al. [13] Our patient also had gastric involvement with a clinical suspicion of malignancy at the time of surgery. On gross pathological examination, the ulcers are the most common manifestation, often large, with rolled and irregular edges that may mimic malignancy as was suspected in our case too. Disseminated zygomycosis rapidly invades vessels resulting in thrombus formation, with infarction, necrosis, hemorrhage, and perforative peritonitis, which can sometimes mimic a primary ischemic pathology especially in older patients [14] as was seen in our case. There could be spread to multiple organs, most frequently lung, however in our case, there was no evidence suggesting any other organ involvement. [15] The gold standard for diagnosis of zygomycosis is demonstration of tissue invasion by fungal hyphae with a classical morphology in histopathological specimen, which was present in our case. Serologic tests presently have no significant role in diagnosis Zygomycesis and positive fungal cultures are seen in only 50% cases. Polymerase chain reaction can be useful for the diagnosis and for species identification, however is not completely reliable for confirmation of diagnosis. [12] In our case, though culture was advised but patient lost for further follow up.

Histoplasmosis is a systemic fungal infection caused by Histoplasma capsulatum which occurs endemically in some parts of the world but is uncommon in India, with only sporadic reports from different parts of country. [16] Disseminated Histoplasmosis (DH) is an extremely rare mycotic infection, usually occurs secondary to high inoculum infections or direct invasion due to trauma. [17] About 40% of patients do not have any obvious risk factors. In our case too, there were no findings suggesting any definite immunosuppression, except a previous history of irregular AKT treatment. Doleschal et al reported first case in a immunocompetent patient in Austria from a non-endemic area. [18] From India, Baudhha et al have also reported a case of progressive DH in an immunocompetent male from non-endemic area. [19] However, Subramanian et al, in an Indian study analysed 19 cases of DH (10 year period) with three cases of GI involvement and found diabetes mellitus and HIV infection to be most common co-morbid conditions. [20]

In clear contrast to the initial presentation of acute histoplasmosis, which almost exclusively involves the respiratory tract, DH most prominently affects the gastrointestinal tract with the typical sites of involvement being terminal ileum and caecum, as seen in our case.
Majority of the patients have a subclinical disease course [18], and are usually asymptomatic. However in symptomatic patients, the most common presentation is fever, bloody diarrhea, abdominal pain, hepatosplenomegaly and lymphadenopathy. [5, 20] In our case too, abdominal pain, lymphadenopathy and possibly hepato-splenomegaly (abdominal distension) were present. However our patient primarily presented with obstructive features (vomiting, constipation). The clinical presentation including endoscopy and radiology sometime can be mistaken for infections like mycobacteria, Entamoeba histolytica; or inflammatory bowel disease, appendicitis and even malignancy. [17,18] In our case too, as patients had obstructive features associated with a caecal mass on CT, hence clinically, a possibility of malignancy was suggested. Sehgal et al have also reported a case of GIH, who was suspected to have colonic cancer in view of the findings seen on scopy and radiology. [14] Microscopic findings usually seen in GIH are diffuse, often nodular lymphohistiocytic infiltrates, showing fungal elements within the cytoplasm of macrophages are pathognomonic of histoplasmosis, while well-formed granulomas are rare. [19] In our case however, multiple well-formed granulomas were seen.

In tuberculosis-endemic regions as in the Indian subcontinent, DH can easily be mistaken for tuberculosis owing to its similar clinical presentation like prolonged fever, hepatosplenomegaly, lymphadenopathy; and histoplasmosis must always be considered in the differential diagnosis, particularly if there is no response to empirical anti-tubercular therapy (ATT). [21] The Chest X ray findings in DH can also mimic TB as both can show cavitatory lesions and fibrosis. [19] Histology can also be confusing sometimes due to presence of granulomas, and even caseous necrosis. Hence it becomes imperative to use appropriate staining and culture techniques to identify this organism especially in the immunocompromised. [20] In our case too, though there was a possible past history with a clinical suspicion of TB, which however could not be confirmed, neither on FNAC (which rather showed histoplasmosis), nor on culture as that was also not done. Though the patient took irregular AKT treatment for a short period but stopped; either due to poor compliance or due to a possible lack of response due to him being a case of histoplasmosis.

**Conclusion**

GI fungal infections are uncommon, but not rare. They often get missed due to their overlapping clinical features with various other conditions. Fungal etiology can often be mistaken for ischemic pathology owing to angio-invasion and thrombosis; and also sometimes for malignancy leading to extensive surgeries. Hence, a high index of suspicion is most essential, both by clinician and pathologist to make a correct diagnosis. Therefore, a fungal etiology should always be suspected in every case of a GI ulcer or perforation, and pathologist should always ask relevant special stains. Moreover, a simple, but timely treatment with appropriate antifungal therapy is very crucial as the mortality is still high with certain fungi, especially in patients with immunosuppression, including old age and diabetes.

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**References**


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