Segmental dilatation of ileum in a young patient presenting with anemia

Neha Nischal, Deepak Balachandra, Anil Kumar Agarwal, Sunil Kumar Puri
Departments of Radiology and Gastro-intestinal Surgery, GB Pant Institute of Postgraduate Medical Education and Research (GIPMER), New Delhi, India

Correspondence: Dr. Neha Nischal, Department of Radiology, GB Pant Hospital, 1, JLN Marg, New Delhi - 110 002, India.
E-mail: neha.nischal@gmail.com

Abstract
Segmental dilatation of ileum (SDI) is a less known uncommon entity with a confusing clinical scenario and no definite etiopathogenesis. The preoperative diagnosis is of exclusion. However, it has an excellent prognosis after surgery. We describe a case of a young patient who presented with anemia without any overt gastrointestinal (GI) bleed. Thorough radiological examinations were needed to reach the diagnosis of SDI which was confirmed postoperatively.

Key words: Barium studies; ileal dilatation; segmental dilatation

Introduction
Segmental dilatation of ileum (SDI) is a rare entity characterized by an idiopathic sharply demarcated segment of dilated ileum. It is difficult to diagnose clinically in view of absence of specific signs and symptoms. We describe the case of a young patient with SDI with emphasis on the clinico-radiological aspects of this entity.

Case Report
A 14-year-old male patient presented with complaints of vague abdominal pain and fullness on the right side of abdomen. He also had history of intermittent episodes of low-grade fever since 4 months. Clinical examination revealed mild distension on the right side of abdomen without a palpable mass. Laboratory investigations were normal except for anemia with Hb of 6.8 g% (normal range 13.5–17 g% for males). However, there was no history of any overt gastro-intestinal (GI) bleeding. Abdominal radiograph showed a large gas shadow in the right hypochondrium along with few enteroliths on the right side of abdomen [Figure 1]. An ultrasound of the abdomen revealed a large unilocular cystic lesion showing air–fluid level in the right lumbar and iliac region. Contrast-enhanced computed tomography (CECT) of the abdomen revealed similar imaging findings with the cyst showing air–fluid level and appearing to be in continuity with the ileum, suggesting dilated ileal loop [Figures 2 and 3]. However, no cause for obstruction could be found on CT. Barium meal follow-through (BMFT) examination was carried out later which proved that the cystic lesion was an aneurysmally dilated ileal loop with normal caliber of ileum proximal and distal to the dilated segment. The dilated segment showed normal peristalsis without evidence of any obstruction and was oriented along the long axis of the ileum [Figures 4 and 5].

Intraoperatively, there was large cystic dilatation of the mid-ileum, measuring about 15 × 14 cm, located about 90 cm from the ileocecal valve. The cyst was excised en-bloc along with 30 cm of the ileum and the jejunum was anastomosed to the cecum. The postoperative period was uneventful, and the patient was discharged on the 7th postoperative day. The histopathological examination of the resected specimen showed a cystic dilatation of ileum without any evidence of ulceration or neoplasia.

This is an open access journal, and articles are distributed under the terms of the Creative Commons Attribution-NonCommercial-ShareAlike 4.0 License, which allows others to remix, tweak, and build upon the work non-commercially, as long as appropriate credit is given and the new creations are licensed under the identical terms.

For reprints contact: reprints@medknow.com

Cite this article as: Nischal N, Balachandra D, Agarwal AK, Puri SK. Segmental dilatation of ileum in a young patient presenting with anemia. Indian J Radiol Imaging 2018;28:369-72.
proximal to the ileo-caecal junction causing postero-lateral displacement of the ascending colon [Figure 6]. The cut section showed normal mucosal folds in the dilated segment with $3 \times 2$ cm irregular mucosa. Segmental resection with end-to-end anastomosis of the involved ileum was done. Postoperative course was uneventful. Biopsy showed normal bowel musculature and ganglion cells. An island of ectopic gastric mucosa was found corresponding to the irregular mucosa on gross examination. Final diagnosis of segmental dilatation of ileum (SDI) was made.

### Discussion

SDI, also known as segmental mega-ileum or ileal dysgenesis, is an unusual congenital dilatation of a segment of bowel which most commonly involves ileum.[1] The entity was initially reported by Swenson and Rathauzer in 1959. Their case involved the colon. Since then, fewer than 200 cases have been published worldwide.[2] They also proposed the following criteria for diagnosing this entity: (1) limited dilatation of bowel with threefold or fourfold increased size; (2) sharp conversion between dilated and normal intestine without an obvious obstruction; (3) the
presence of normal neuronal plexus in submucosa and muscularis propria; (4) clinical presentation of obstruction or sub-obstruction; and (5) recovery after resection of the dilated segment.\[^3,4\]

Presentation is commonly in infancy or childhood and symptoms vary with age at presentation. In the neonatal period, obstruction is the commonest feature owing to the aperistaltic bowel segment. However, later in life, the patient may present with GI bleeding, anemia, abdominal pain, or may be asymptomatic.\[^2,4,5\] The cause of overt or occult GI bleeding is attributed to the presence of ulcers and/or ectopic gastric mucosa within the dilated segment. A high proportion of these cases are reported to have other GI tract anomalies including omphalocele, malrotation, and Meckel’s diverticulum.\[^1-4,6,7\] A case of SDI with anterior thoracolumbar meningocoeele and bilateral undescended testes has also been described in literature.\[^9\] In this report, the patient had vague abdominal pain with asymptomatic anemia, without any associated GI anomalies.

The etiopathogenesis is still debatable. However, most authors postulate an extrinsic source of compression in

Radiological examinations are quite contributory to the diagnosis. Abdominal radiograph usually reveals dilated bowel loop on the right side with or without air–fluid levels, which may suggest possibility of cecal volvulus in the setting of intestinal obstruction. Ultrasound reveals a large cystic lesion, which may mimic a mesenteric cyst or duplication cyst of the small bowel. Barium studies disclose the nature of the lesion as being a dilated segment along the long axis of bowel, as opposed to Meckel’s diverticulum or other small bowel diverticuli which show anti-mesenteric or mesenteric orientation. Other differentials may include aneurysmal dilatation in lymphoma or cavitating neoplasms, but CT usually suffices in ruling out these causes.\[^3,6,8\] CT enteroclysis is reported to be even more sensitive in diagnosing this lesion accurately by achieving adequate distension of bowel.\[^6\]

Exclusion of Hirschsprung disease is imperative before making the diagnosis of SDI where there is normal neuronal plexus within the dilated segment.\[^3\]

Many authors have also labeled this entity as a giant Meckel’s diverticulum. However, there does exist some differences between the two, most important being the anti-mesenteric orientation of the Meckel’s diverticulum as opposed to SDI, which is oriented along the long axis of bowel. Heterotopic gastric or pancreatic mucosa may be found in both the entities and should not be the criteria for diagnosing giant Meckel’s diverticulum.\[^9,30\]
The treatment is resection of the involved segment with end-to-end bowel anastomosis which has an extremely favorable outcome for the patient.[1-5,7]

Conclusion

In the light of pauci-clinical manifestations, imaging has paramount role in reaching the correct preoperative diagnosis of SDI. High index of suspicion should be kept in young patients with unexplained anemia or GI bleed showing cystic dilatation of ileum. We also propose to change the nomenclature of giant Meckel’s diverticulum because of the inherent differences between the two entities.

Declaration of patient consent

The authors certify that they have obtained all appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

Financial support and sponsorship

Nil.

Conflicts of Interest

There are no conflicts of interest.

References

1. Sam CJ. Segmental ileal dilatation in a child. Trop Gastroenterol 2011;32:221-3.
2. Paradiso FV, Coletta R, Olivieri C, Briganti V, Oriolo L, Fabbri R, et al. Antenatal ultrasonographic features associated with segmental small bowel dilatation: An unusual neonatal condition mimicking congenital small bowel obstruction. Peditr Neonatol 2013;54:339-43.
3. Wei CH, Sheu JC. Concomitant segmental intestinal dilatation and omphalocele. Formos J Surg 2011;44:168-70.
4. Khemakhem R, Riazulhaq M, Elhassan EO. Segmental dilatation of intestine presenting as partial intestinal obstruction in a child. APSP J Case Rep 2014;5:19.
5. Sjolin S, Thoren L. Segmental dilatation of the small intestine. Arch Dis Child 1962;37:422-4.
6. Lee RKL, Hung EHY, Leung JHY, Tsang KWK. Idiopathic localised dilatation of the ileum: Computed tomography enteroclysis. Hong Kong J Radiol 2014;17:124-8.
7. Raj P, Sarin YK. Segmental dilatation of ileum associated with anterior thoracolumbar meningomyelecele and bilateral undescended testes. APSP J Case Rep 2015;6:5.
8. Javors  BR, Gold  RP, Ghahremani GG, Radin DR, Cho KC, Maglinte DD, et al. Idiopathic localized dilatation of the ileum in adults: Findings on barium studies. Am J Roentgenol 1995;164:87-90.
9. Akbulut S, Yagmur Y. Giant Meckel’s diverticulum: An exceptional cause of intestinal obstruction. World J Gastrointest Surg 2014;6:47-50.
10. Nunes Q, Hotouras A, Tiwari S, Sheth A. Gangrene due to axial torsion of a giant Meckel’s diverticulum containing multiple stones in the lumen: A case report. Cases J 2009;2:7141.