An Atypical Presentation of Giant Meckel’s Diverticulitis: A Case Report

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Abstract

Meckel’s Diverticulum (MD) is the most common congenital malformation of the gastrointestinal tract and the only true diverticulum of the small bowel, resulting from the incomplete obliteration of the vitelline duct in the first 5 to 7 weeks of gestation. However, MD is rare, with a traditional prevalence of 2% in the general population. While the majority of MD never becomes symptomatic, potential for severe complications may arise secondary to diverticulitis with or without perforation, hemorrhage and obstruction. Considerable debate therefore exists whether or not to surgically resect MD found incidentally. Moreover, such complications present considerable diagnostic challenges, and given its rarity, are scarcely considered in the differentials of an acute abdomen.

We present one of the longest cases of giant MD reported in the literature, with non-perforated diverticulitis in a young adult male attending with an acute abdomen and normal inflammatory markers. He underwent successful, un-complicated laparoscopic resection. Histopathological analysis confirmed Meckel’s diverticulitis in the absence of ectopic gastric or pancreatic tissue.

Introduction

Meckel’s Diverticulum (MD) is the most common congenital abnormality of the gastrointestinal tract [1,2]. First reported by Johann Meckel, it arises from the incomplete involution of the omphalomesenteric duct in the 7th week of gestation [1,3,4], and is the only true diverticulum of the small intestine [5].

The maxim of twos is well established in traditional surgical teaching of MD: typically 2 inches long, 2 feet proximal to the ileocaecal valve, present in 2% of the population, twice as common in males and most often symptomatic in patients aged less than 2 years. Recent evidence has shown prevalence between 0.3% to 2.9% of the population [6-13]. Moreover, the largest systematic review of recent literature calculated a weighted mean distance of 52.3cm (7 to 200) from the ileocaecal junction on the antimesenteric border of the ileum, and a mean weighted length of 3.05 cm (0.4 to 11), approximately 1 inch, with a mean diameter of 1.58 cm (0.3 to 7) [2].

Giant MD, considered greater than 2 inches (or 5 cm), considerably increases the risk of more severe forms of complications, particularly intestinal obstruction [14,15]. Increasing evidence also suggests a further correlation between size and severity of symptoms [2,16].

To our knowledge, herein we present one of the largest cases of giant MD reported in the literature, with implications for definitive surgical management.

Case Presentation

A 23-year old Caucasian male presented to the acute surgical take with a short history (several hours) of generalized abdominal pain and one episode of vomiting. Past medical history included laparoscopic appendicectomy in 2015 with histology revealing a neuroendocrine tumour. The patient was subsequently lost to follow-up. He was otherwise fit and well, on no regular medications and with no known allergies. On examination his observations were within normal limits and he was maximally tender in the right iliac fossa. Blood results were borderline normal, with a mild leukocytosis (white cell count 12.0 UNITS) and C-Reactive Protein (CRP) <3 UNITS. Urea and electrolytes, liver function tests and serum amylase were within the normal range. Plain radiographs of the chest and abdomen were unremarkable.
Given his past history of malignancy and clinical findings, a Computed Tomography (CT) scan was performed (Figure 1), which showed a fluid-filled mass extending from the small bowel into the pelvis consistent with a Meckel’s diverticulum. Diagnostic laparoscopy showed a MD 15.24 cm in length (Figure 2), which was excised using a laparoscopic surgical stapling device (Endo GIATM). The specimen was sent for histological analysis. The patient made a slow recovery jaded by a post-operative ileus which resolved conservatively and he was subsequently discharged home. Histological findings indicated small intestinal type villous mucosa with no evidence of gastric or pancreatic mucosa, with severe inflammation and ulceration in keeping with Meckel’s diverticulitis. No evidence of neoplasia was found.

**Discussion**

MD is the only true diverticulum of the GI tract, comprising all three layers of the bowel wall and results from the incomplete atrophy of the vitelline duct. While 90% of MD are between 1 cm to 10 cm in length, ‘giant MD’ exceeding 5 cm are rare [17], with the largest recorded nearly a century ago measuring over 100 cm [18].

Although the majority of MD remain asymptomatic and are discovered incidentally, the lifetime incidence of symptomatic presentation remains between 4% to 9% [4,9,19], compared to 7% to 8% in appendicitis [16].

Intestinal obstruction is the most common complication of symptomatic MD, followed by hemorrhage from peptic ulceration with heterotopic mucosa, and diverticulitis with or without perforation, as in this case [20], accounting for 90% of symptomatic MD [19]. Obstruction arises from multiple aetiologies, including intussusceptions, whereby the MD is the lead point; mechanical volvulus around a persistent fibrous band attaching the MD to the umbilicus, or axial twisting around a narrow base; diverticular stricture; Littre’s herniation, and inflammatory adhesions [2]. Rarer complications include umbilical abnormalities such as fistulation and Meckelian cancers, generally diagnosed after 60 years of age [21,22].

Risk factors for developing complications are well established. Male sex, age under 50 years, length greater than 2 cm, and heterotopic presence of gastric or pancreatic mucosa increase the risk of complicated MD [23]. Presence of two, three (seen here) or four criteria increases the symptomatic prevalence to 25%, 42% and 70% respectively [23]. Nonetheless, a high index of suspicion is necessary for prompt diagnosis and treatment [24], with only 4% of MD identified pre-operatively either clinically or radiologically [25]. The presence of heterotopic mucosa (gastric and pancreatic tissue accounting for 97%) is the most significant factor for determining the need for surgical intervention, and is closely correlated with hemorrhage. Symptomatic MD decreases with age [10,13,26-30], with over half of all children with MD requiring surgery under 5 years [31]. Interestingly, Negrea V et al. [32] found a higher nerve fiber density in the walls of the Meckel’s lined with intestinal mucosa, as seen in this case, compared to areas lined with ectopic gastric mucosa and the walls of the ileum [32]. An inverse correlation exists between nerve fiber density and age, with higher nerve fiber density resulting in more intense local peristalsis, which may explain the observed predisposition to intussusception [32].

Giant MD has a higher incidence of obstruction [19,33]. Moreover, diverticulitis, torsion and volvulus are more common complications in longer MDs with a narrow base, while short MDs with a wider base stand at higher risk of intussusceptions [34]. Thus, an elongated variant with a narrow neck is more likely to result in torsion and diverticulitis, as seen in this case, whereas short, wide-base diverticula may promote foreign body entrapment [15]. To the author’s knowledge, this is one of the longest MD reported in the literature, with an abnormal presentation with near normal inflammatory markers.

Definitive management remains surgical excision of the MD with
or without resection of adjacent small bowel, the latter preferred in the presence of severe inflammation [34]. Laparoscopic procedures have been shown to be safe and confer no worse outcomes or complications than open surgery, through single or 3 trocars, intra-peritoneally or exteriorization [35,36]. Debate exists whether silent MD should be resected when incidentally discovered. Recent reviews by Zani et al. [9] and Soltero et al. [19] found that resection of silent MD conferred a significantly higher post-operative complication rate than leaving in situ. Zani et al. [9] found a 5.3% risk of postoperative complications after prophylactic resection and a 1.3% risk of developing symptoms after leaving it in situ. They also found no long-term complications associated with leaving the Meckel's in situ when reviewing articles that reported follow-up on patients with silent Meckel's left in situ, and estimated that more than 750 silent MD would have to be resected in order to preserve one life.

**Conclusion**

This case demonstrates the highly variable presentation of giant Meckel's diverticulitis, conferring considerable diagnostic and, given its considerable length, operative challenges. A diagnosis of MD should always be considered in the differential of an acute abdomen presenting with right iliac fossa pain, and multi-modality investigations including CT are recommended.

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**References**


