Meckel’s diverticulum — a case study

Sandhyarani M. PATIL[1]+
Mahadeo Ramchandra PATIL[2]

Department of Anatomy, Rajarshi Chhatrapati Shahu Maharaj Government Medical College [1], Department of Surgery, Dr. D. Y. Patil Medical College [2], Kolhapur, INDIA.

+ Dr. Sandhyarani M. Patil, MS  
Assistant Professor  
Department of Anatomy  
Rajarshi Chhatrapati Shahu  
Maharaj Govt. Medical College  
Kolhapur, INDIA.  
☎ +91 982 2286190 
✉ drsandhyapatil.2009@gmail.com

Key words [Meckel’s diverticulum] [congenital] [gastrointestinal tract]

Abstract
This paper presents and describes a case of Meckel's diverticulum. It was found in a 10-year-old boy who was operated for appendectomy in surgery department. It was located 24 cm proximal to the ileocaecal junction. It was 4.3 cm in length with broad tip and 3 cm in diameter. It was attached to antimesentric border. The tip was not attached by any fibrous band to umbilicus.


Introduction
The Meckel's diverticulum or diverticulum ilei is the most common unusual congenital condition of gastrointestinal tract. It is present in fetal life as the vitello-intestinal duct, which usually disappears later in seventh week of gestational period.

The incidence of Meckel's diverticulum in the general population has been estimated to be about 2% [1]. Reports from autopsy and retrospective studies range from 0.14-4.5% [2, 3].

Although Meckel's diverticulum occurs in both sexes, it may cause complications more frequently in males, and therefore is often diagnosed in males [4–6].

Ninety per cent of diverticula are within 90 cm of distance to the ileocaecal valve, although diverticula up to 180 cm from ileocaecal valve have been observed. A person with Meckel's diverticulum has 4-6% lifetime risk of developing a complication [7]. The major complications are diverticulitis and perforation [6].

Case Report
Ten-year-old boy was operated for appendectomy in the Department of Surgery at Chhatrapati Shahu Maharaj Goverment Medical College. On examination of bowel, Meckel's diverticulum was found. It was removed and the specimen of Meckel's diverticulum was processed for histological study, where it was stained with hematoxylin and eosin. Following features were noted: the Meckel's diverticulum was seen at 22 cm proximal to ileocaecal junction; it was 4.3 cm in length with broad tip; it was 3 cm in diameter; it was attached to antimesentric border; its tip had no attachment via fibrous band to umbilicus (Figure 1).

Histology
The mucus membrane showed villi. The villi were lined by simple columnar epithelium with goblet cells. The lamina propria showed glands along with isolated lymphatic nodule. Submucosa was present. Muscle layer consisted of inner circular and outer longitudinal layer and outermost serosa (Figure 2).

Discussion
Developmentally the vitelline duct connects the yolk sac and midgut of the embryo at an early stage in fetal life for nutrition [5]. Thus, in the 3 week old embryo the yolk forms the ventral aspect of the gut and communicates through a wide, short vitelline duct, which subsequently becomes absorbed. The blood supply to the vitelline duct comes from paired ventral branches of the abdominal aorta, the vitelline arteries.

If this duct does not involute by 7th week of gestation, various developmental remnants may be left. Failure of most proximal
portion of duct to obliterate results in a Meckel’s diverticulum [8]. The yolk sac is continuous with the embryo and contains all layers of the intestinal wall [8].

Meckel’s diverticulum may remain completely asymptomatic, or it may mimic disorders such as Crohn’s disease, appendicitis and peptic ulcer disease. Ectopic tissue found in approximately 50% of cases consists of gastric tissue in 60-65% of cases and pancreatic tissue in 5-16% [9].

Meckel’s diverticulum occurs more commonly in individuals born with other congenital malformations especially exomphalos, esophageal atresia, anorectal atresia and gross malformations of the cardiovascular and central nervous system [9]. Hemingway and Allisan reported an association between angiodysplasia of the caecum and ascending colon with Meckel’s diverticulum in five patients aged between 13 and 21 years [10].

Knowledge of Meckel’s diverticulum is important for surgeons to avoid complications during various abdominal surgeries. It is also important for radiologists while doing ultrasound examination and evaluating radiographs.

References