Case Report

A Long Rectal Duplication Cyst in an Adult Cadaver: A Rare Anatomical Variation

Aneesh Dave*, Rohan Dalal, Bardia Aryaie and Bahadir Cem Demirdes
Department of Anatomy and Histology, The University of Sydney, Australia

Abstract
This report describes a long rectal duplication cyst found during routine dissection of an 82 year old male cadaver. The cyst was tubular, blind-ending and non-communicating. It emerged from the posterior rectal wall approximately 13 cm superior to the anal verge and was 12 cm in length. Long duplication cysts such as that described may cause complications such as perforation, bleeding and chronic pain. Hence duplication cysts should form part of the differential work up in a clinical setting.

Introduction
Duplication cysts are a congenital malformation which may develop along any part of the alimentary tract, occurring with an estimated prevalence of 1 in 4500 cases [1]. Duplication cysts involving the rectum are very rare, accounting for only 5% of all alimentary tract duplications [2]. Of these, 80% are spherically shaped, and 80% of patients present before two years of age [3]. Only a small number of case reports describe rectal duplication cysts presenting in adults. This case report describes a tubular, blind-ending rectal duplication cyst found during a routine dissection of an 82 year old male cadaver. Several theories have been developed to explain the embryological development of rectal duplication cysts; however none of these have been conclusive [4]. Rectal duplication cysts are an important clinical consideration, as they may result in obstruction, perforation and chronic pain [5] and must therefore be considered within a differential diagnosis framework.

Case Report
Gross pelvic dissection of an 82-year old male cadaver of European ancestry was performed as per a recognized and widely used dissection manual [6], whereby a variant long rectal duplication cyst was noted. Incidentally, this cadaver was also noted to have an aberrant inferior mesenteric artery arising from the coeliac trunk. Otherwise, in life there were no significant medical issues, and there was no history of any surgical procedures to the area.

The opening of the cyst was found on the posterior aspect of the rectum, approximately 135 mm superior to the anal verge (Figure 1). The duplication cyst was noted to be 124 mm in length.

Abstract
This report describes a long rectal duplication cyst found during routine dissection of an 82 year old male cadaver. The cyst was tubular, blind-ending and non-communicating. It emerged from the posterior rectal wall approximately 13 cm superior to the anal verge and was 12 cm in length. Long duplication cysts such as that described may cause complications such as perforation, bleeding and chronic pain. Hence duplication cysts should form part of the differential work up in a clinical setting.

Introduction
Duplication cysts are a congenital malformation which may develop along any part of the alimentary tract, occurring with an estimated prevalence of 1 in 4500 cases [1]. Duplication cysts involving the rectum are very rare, accounting for only 5% of all alimentary tract duplications [2]. Of these, 80% are spherically shaped, and 80% of patients present before two years of age [3]. Only a small number of case reports describe rectal duplication cysts presenting in adults. This case report describes a tubular, blind-ending rectal duplication cyst found during a routine dissection of an 82 year old male cadaver. Several theories have been developed to explain the embryological development of rectal duplication cysts; however none of these have been conclusive [4]. Rectal duplication cysts are an important clinical consideration, as they may result in obstruction, perforation and chronic pain [5] and must therefore be considered within a differential diagnosis framework.

Case Report
Gross pelvic dissection of an 82-year old male cadaver of European ancestry was performed as per a recognized and widely used dissection manual [6], whereby a variant long rectal duplication cyst was noted. Incidentally, this cadaver was also noted to have an aberrant inferior mesenteric artery arising from the coeliac trunk. Otherwise, in life there were no significant medical issues, and there was no history of any surgical procedures to the area.

The opening of the cyst was found on the posterior aspect of the rectum, approximately 135 mm superior to the anal verge (Figure 1). The duplication cyst was noted to be 124 mm in length.

Abstract
This report describes a long rectal duplication cyst found during routine dissection of an 82 year old male cadaver. The cyst was tubular, blind-ending and non-communicating. It emerged from the posterior rectal wall approximately 13 cm superior to the anal verge and was 12 cm in length. Long duplication cysts such as that described may cause complications such as perforation, bleeding and chronic pain. Hence duplication cysts should form part of the differential work up in a clinical setting.

Introduction
Duplication cysts are a congenital malformation which may develop along any part of the alimentary tract, occurring with an estimated prevalence of 1 in 4500 cases [1]. Duplication cysts involving the rectum are very rare, accounting for only 5% of all alimentary tract duplications [2]. Of these, 80% are spherically shaped, and 80% of patients present before two years of age [3]. Only a small number of case reports describe rectal duplication cysts presenting in adults. This case report describes a tubular, blind-ending rectal duplication cyst found during a routine dissection of an 82 year old male cadaver. Several theories have been developed to explain the embryological development of rectal duplication cysts; however none of these have been conclusive [4]. Rectal duplication cysts are an important clinical consideration, as they may result in obstruction, perforation and chronic pain [5] and must therefore be considered within a differential diagnosis framework.

Case Report
Gross pelvic dissection of an 82-year old male cadaver of European ancestry was performed as per a recognized and widely used dissection manual [6], whereby a variant long rectal duplication cyst was noted. Incidentally, this cadaver was also noted to have an aberrant inferior mesenteric artery arising from the coeliac trunk. Otherwise, in life there were no significant medical issues, and there was no history of any surgical procedures to the area.

The opening of the cyst was found on the posterior aspect of the rectum, approximately 135 mm superior to the anal verge (Figure 1). The duplication cyst was noted to be 124 mm in length.
and 11.8mm in diameter. During its course, it curved around the rectosigmoid junction passing anterior to the rectum. The duplication cyst was a blind-ending, non-communicating tubular-shaped cyst. As the duplication cyst was laid open, it was noted to have a thick wall, corresponding to the multi-layered wall consistent with normal rectal lining including a mucosa and muscular layer (Figure 2). This allowed differentiation between a true duplication cyst and a simple rectal diverticulum.

No other regional anatomical variants were noted. There was no significant past history of pelvic surgery or pelvic pathology associated with the cadaver.

Discussion

Duplication cysts involving the rectum are a rare congenital malformation, accounting for only 5% of all duplications in the gastrointestinal tract. Of those, the majority of cysts are reported as being within the retro-rectal space, spherically shaped, presenting in children less than two years of age and with predominance in males of European ancestry [3,7-9]. Presentation of rectal duplication in adults, as described in this case study, has only been described within a European ancestry [3,7-9].

As the duplication cyst was laid open, it was noted to have a thick wall, corresponding to the multi-layered wall consistent with normal rectal lining including a mucosa and muscular layer (Figure 2). This allowed differentiation between a true duplication cyst and a simple rectal diverticulum.

The mechanism of development of rectal duplications is not fully understood, with several theories attempting to explain their formation. The mostly widely recognized theory involves an abnormality during the gastrulation stage resulting in a split notochord [4]. The herniation of the yolk sac through a gap between the ectoderm and endoderm eventually is thought to result in a yolk sac to amniotic cavity fistula, which is closed to varying degrees through continued growth of the embryo. Although this can cause duplications of the entire gastrointestinal tract, not all duplication cysts can be explained with this theory.

The presence of rectal duplication cysts should not be missed, as there is a possibility of serious complications including obstruction, intussusception, infection and volvulus which may be associated with duplication cysts. The presence of ectopic gastric mucosa, which occurs in approximately a third of rectal duplication cysts [2], can also cause perforation and haemorrhage. Even if such complications are not present, constipation, chronic pain and abdominal distension can be a cause of significant morbidity to the patient [5]. In a case series of 72 patients by Stringer et al., it was noted that spinal and genitourinary anomalies were present in two of six patients, and thus further evaluation may be warranted in patients with noted rectal duplication cysts [9]. Therefore, due to the highly variable, non-specific presentation, the high risk of associated co-morbidities and the difficulties in diagnostics, rectal duplication should be considered as a differential diagnosis in the refractory patient.

Acknowledgement

The authors would like to acknowledge Dinuke De Silva and Steven Gluckman for assisting with dissection.

References


Figure 2: Internal features of rectal duplication cyst.
A - Rectum, B - Opening of rectal duplication cyst; C - Mucosal Lining of cyst.