Case report

Rare benign tumours of the small intestine presented as acute abdomen in childhood

B. Kasselas, M. Papoutsakis, Ch. Chaidos, Ch. Kasselas, G. Tsikopoulos, M. Agelidou, G. Papouis, Th. Karagiozoglou

SUMMARY

Benign tumours of the small intestine in children although rare, may represent an additional cause of acute abdomen symptoms that the gastroenterologist, the paediatrician and the surgeon must keep in mind in order to proceed to accurate diagnosis and appropriate treatment. Three cases of benign tumours of the small intestine, histologically diagnosed as leiomyoma, fibroma and hamartoma, are presented, and the literature is revised emphasizing issues of pathophysiology, specific characteristics and the treatment of these tumours.

Key words: Benign tumours, small intestine, acute abdomen, children.

INTRODUCTION

Neoplasms of the small intestine represent a percentage of 3% to 6% of the tumours of the gastrointestinal tract and less than 2% of all tumours.\(^1,2\) Benign tumours of the small intestine are estimated to be 10% of all non-malignant tumours of the gastrointestinal tract and 30% of all the neoplasms of the small intestine.\(^3,4,5\) Among benign neoplasms of the small intestine leiomyoma, adenoma, lipoma, haemangioma, fibroma and hamartoma are included with regard of their incidence.\(^2,6,7,8\) These neoplasms are usually seen after the fifth decade of life with a peak incidence between 70 to 80 years of age.\(^7\)

CASE NR1

A 4 year old boy presented complaining of abdominal pain, bilious vomiting, and fever up to 38.5 degrees. The symptoms of the disease started two days before presentation and no previous history was reported. The diagnosis of acute abdomen was established by clinical findings, physical examination, diagnostic tests, radiography and laboratory investigations (leucocytosis with neutophilia, radiological signs of ileus of the small bowel). During laparotomy a tumour of the small bowel was revealed at the mesenteric margin of the small intestine (approximately in the middle of the bowel). It was of a hard consistency with a diameter of 1.2cm, limited by the surrounding tissues; its colour was dark red and it provoked a large subserosal haematoma measuring 5 cm on both sides (Fig. 1). Because of all these characteristics, the tumour was considered benign. The exploration of the rest of the abdomen indicated no other

Figure 1. Macroscopic appearance of the leiomyoma

Pediatric Surgery Dept, Hippokration Hospital, Thessaloniki, Greece

Author for correspondence:
Th. Karagiozoglou, Ethnikis Amyntis 14, 546 21 Thessaloniki, Tel.: +2310.265.724, e-mail: thomaiskl@hotmail.com
pathological signs and an intestinal resection of 5cm on both sides of the tumour (with excision of the mesentery) with an end to end anastomosis was performed. The postoperative course of the young patient was without complications. A leiomyoma of the small bowel with no histological evidence of malignant characterisitics was diagnosed (Fig 3) which was completely excised and had no regional lymph node metastasis. To date, 7 years after the operation, the patient has had no other problems.

CASE NR2

An 11 year old girl was hospitalized in the pediatric dept. for anemia investigation. The disease manifested with diffuse, recurrent, non-typical abdominal pain, anorexia, vomiting, constipation and weight loss (7kg.) over the previous six months. Profound anemia was evidenced through physical examination and diagnostic tests (Hb 6 gr. Ht 19%). Further laboratory investigations were normal except for a positive Mayer stool test. Screening tests and radiological investigation provide no information, as did the scintigram (Tc-99) which was normal. On the other hand, the ultrasound examination revealed a small fluid collection in Douglas space. The anemia was treated with blood transfusion and a CT study of the abdomen was scheduled. During the investigation, the patient showed signs of high obstructive ileus so she was subjected to an emergency operation. Exploratory laparotomy revealed the existence of a jejuno-jejunal intussusception. After manual reduction, a tumour was palpated (as the lead-point causing intussusception) whitish in color, of hard consistency, with a diameter of 1.5cm and which was localized at the antimesenteric margin of the bowel and occupying almost the whole intestinal lumen (Fig 2). Segmental resection of the small intestine with tumour free margins was performed as well as an end-to-end anastomosis. Surgical exploration of the abdominal cavity failed to reveal any additional findings. The postoperative course of the patient was uncomplicated. Histology diagnosed an intestinal fibroma with no malignant evidence (Fig 4), whereas the margins of resection and the regional lymph nodes were not infiltrated. To date, three years later, the patient, who is examined periodically, has had no other health problems.

Figure 2. Macroscopic appearance of the fibroma

Figure 3. Microscopic appearance of the leiomyoma

Figure 4. Microscopic appearance of the fibroma
CASE NR3

This case refers to a 6 month-old boy, who presented with abdominal pain, vomiting plus blood-stained mucus in the stools for the previous 12 hours. From the history and physical examination, plain X-rays of the abdomen (that showed signs of obstruction) and ultrasound examination, possible ileocolic intussusception was considered, which was confirmed during laparotomy. Manual reduction followed and an intraluminal mass the size of a “nut” (5 cm from the ileocecal valve) was palpated (it was the lead-point of intussusception). Its consistency was rough and was limited by the surrounding tissues. A wedge resection of the lesion was performed, followed by an end-to-end ileo-ileal anastomosis. There were no additional findings during surgical examination of the abdomen. Gross pathological examination of the lesion revealed a dark red polypoid projection of the mucosa. Microscopic histological examination showed that it was a myoepithelial hamartoma of the small intestine with no malignant features, whereas the overlying mucosa showed signs of ischaemic necrosis (Fig 5). The patient’s postoperative course was satisfactory. Today, the patient remains completely healthy.

DISCUSSION

Among all benign neoplasms of the small intestine, leiomyoma represents a percentage of 30-35%, hamartomas 10-12% and fibromas 6%6. Their incidence increases from the duodenum to ileum. Approximately half of them are usually asymptomatic and are verified at autopsy.6,7 The main presenting symptoms are abdominal pain, intestinal obstruction with or without intussusception, gastrointe-
precise diagnosis, even though the distinction between a benign leiomyoma and a differentiated one or a leiomyoblastoma, a benign fibroma, a benign myoepithelial hamartoma and a differentiated one may not be always be obvious. Histological indicators for the determination of tumour malignancy are increased mitotic activity (mitoses more than 10 X 10 HPF), cellular pleomorphism, the degree of cellular differentiation and hypercellularity. 

The third question concerns the patient’s management for complete cure. From the previously mentioned evidence, in accordance with the relevant references, it is clear that the tumour must be excised radically by a segmental intestinal resection and excision of the corresponding mesentry, even if there is no macroscopic evidence of malignancy. In these cases, prognosis is excellent. Moreover, if the microscopic examination confirms the complete excision of the tumour, in addition to the absence of any malignant features, no further treatment is required.

A serious problem is presented when the tumour displays macroscopic features of malignancy, or when there is histological documentation that the margins of resection are not free of tumour cells, as well as the identification of hypercellularity or nontypical mitotic activity.

These cases demand more radical operations or other management, and a thorough follow-up of the patient since they carry a high percentage of regional recurrence and a poor prognosis.

REFERENCES