Case Report

Appendiceal mucocele with pseudomyxoma peritonei mimicking ovarian tumor with peritoneal carcinomatosis

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Abstract

Appendiceal mucocele with unprompted pseudomyxoma peritonei is a rare malignant tumor, which is difficult to diagnose before surgery. We present a case of a 62-year-old woman, and subsequently discuss the clinical and imaging presentation of mucoceles. Findings on CT scan suggested 2 diagnoses: appendiceal mucocele with pseudomyxoma peritonei and malignant ovarian tumor with peritoneal carcinosis. MRI suggested the former diagnosis, showing a large tubular mass at the base of appendix with discontinuous wall, and no ovarian abnormality, which was confirmed by mini laparotomy and histological study of peritoneal thickened tissue. Treatment consisted of neoadjuvant chemotherapy, with a good response. This case shows the role of imaging in the preoperative diagnosis of appendicular mucoceles; CT scan and MRI, which are useful tools in identifying undetermined lesions of the appendix.

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Observation

A 62-year-old woman without significant history, presented with abdominal pain located in the right lower quadrant with progressive abdominal distension.

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Laboratory tests showed elevated tumor markers (ACE = 138.08 ng / ml, CA 125 = 306.60 U arb/ml, CA 19-9 = 527.85 U/ml). CT showed a cystic mass in the right lower quadrant, in contact with the caecum, containing curvilinear mural calcifications, and a pelvic cystic mass, it also showed nodular thickening of the peritoneal reflections with stranding and thickening of the omentum (omentum cake), associated with scalloping of the liver surface (Fig. 1). Suggesting 2 main diagnoses; appendiceal mucocele with pseudomyxoma peritonei and malignant ovarian tumor with peritoneal carcinomatosis.

Subsequently an MRI was performed, showing a cystic mass attached to the caecum, with a discontinuous wall, with curvilinear calcifications. On the other hand, the ovaries had no abnormality (Fig. 2). The more likely diagnosis was therefore an appendiceal mucocele with pseudomyxoma peritonei.

A biopsy of the thickened peritoneal tissue was performed by a mini laparotomy, which confirmed the diagnosis of a mucinous cystadenocarcinoma of digestive origin. Appendix removal was not possible due to the presence of multiple adhesions and carcinomatosis nodules. The treatment consisted of neoadjuvant Folfox-based chemotherapy, with positive outcome.

Discussion

The appendiceal mucocele was considered as a pathological entity by Rokitansky in 1842 and officially named by Feren in 1876. Appendiceal mucoceles are uncommon entities from a variety of different pathological processes, of which only a small fraction is associated with the development of peritoneal pseudomyxoma [4].

Two convergent theories have been proposed to explain the disease [7].

Obstruction theory

Appendiceal mucoceles are caused by the accumulation of mucus upstream of appendiceal lumen stenosis (cecal adenocarcinoma, appendix tumor).

Tumor theory

Tumors are the cause of hypersecretion of mucus in the appendix lumen [7].

The clinical flow of the disease does not have a specific picture. It often flows asymptomatically, in about 50% of cases, it is discovered accidentally during radiographic and endoscopic examinations or at surgery [6]. Signs and symptoms, if present, include pain in the right lower quadrant, changes in bowel habits, anemia, hematochezia, and depending on the location of the appendix, other signs, such as hematuria, may be present. The most frightening complication is pseudomyxoma peritonei. Whether benign or malignant, treatment is difficult and prognosis uncertain: its 5-year survival rate is only 53%–75% [1].

The classification of extra-appendiceal disease is also problematic in the evaluation of appendiceal tumors. Surgeons and pathologists use the term “pseudomyxoma peritonei” to describe mucin in the abdomen. Unfortunately, different authors use this term to designate different histological findings. Some authors use the term pseudomyxoma peritonei to describe the accumulation of mucin in the abdominal cavity, whether or not it contains mucous epithelium, while some authors consider it to mean peritoneal carcinomatosis, other groups argue that a distinction should be made between the low-grade pattern of classic PP and true peritoneal carcinomatosis because these 2 entities are associated with a different prognosis and clinical significance [5].

The preoperative diagnosis of appendiceal mucocele is important to select an appropriate surgical approach to prevent peritoneal dissemination, prevent intraoperative and postoperative complications, and repeated surgery [8].

Ultrasoundography is the first-line diagnostic method for patients with acute abdominal pain. It can be used to differentiate mucocele from acute appendicitis. In the context of acute appendicitis, an appendix outer diameter threshold of 6 mm, and greater than 15 mm indicates the presence of a mucocele with a sensitivity of 83% and a specificity of 92% [6]; it shows a cystic, encapsulated lesion of the right lower quadrant, firmly attached to the caecum [4], oblong or piriform with a large vertical axis, with sharp contours [7]. Internal variable echogenicity is related to mucus density. In some patients, multiple echogenic layers along the dilated appendix produce the appearance of ‘onion skin-like’ circles and are pathognomonic for mucocele [4].

CT is considered to be the most accurate diagnostic method. It can be used to detect mucocle-specific signs with high accuracy [10]. A CT scan makes it possible to objectively the connection between the tumor and the cecum. The tumor is often round, well limited, and sometimes partitioned [7]. Cecal lumen greater than 1.3 cm with cystic dilatation and wall calcification [10]. The density of the contents can vary from that of liquid to that of soft tissues (from 10 to 45 HU) with parietal late and homogenous enhancement after injection of iodine [7].

Visceral scalloping, on the other hand, is a diagnostic sign and distinguishes mucinous from fluid ascites on CT. As mucin producing cells in pseudomyxoma peritonei are poorly adherent, the Douglas and/or retro vesical space, the right and left subphrenic spaces, and the surface of the liver and spleen are the commonest sites involved [4].

Dhage-Itatury and Sugarbaker propose a simple yet thorough evaluation of these patients using the new algorithm. Appendectomy alone is the surgical treatment option for patients with benign mucoceles with negative resection margins and no perforation. These patients do not require long-term follow-up [9]. Patients with perforated mucocele, positive margins, positive cytology, and positive appendix lymph nodes should undergo right colectomy and/or cytoreductive surgery (CRS) and/or heated intraperitoneal chemotherapy (HIPEC) or early postoperative intraperitoneal chemotherapy (EPIC) [9].
Fig. 1 – Abdominal post injection CT scan showing a cystic mass in the right lower quadrant, in contact with the caecum (red arrow) containing mural calcifications (white arrow) (A) and pelvic cystic mass (B) (large white arrow). CT also showed nodular and thickening of peritoneal reflections with stranding and thickening of the omentum (head arrow) and peritoneal effusion (Asterisk). (Color version of figure is available online.)
Conclusions

In conclusion, mucocele of the appendix with pseudomyxoma peritonei is a rare disease and can present with confusing symptoms. Preoperative diagnosis greatly assists in determining appropriate management and minimizing both intraoperative and postoperative complications. Ultrasonography, and particularly CT, should be used extensively for this purpose, and sometimes abdominal MRI is a useful tool in identifying undetermined lesions of the appendix.

Patient consent

I confirm that patient has given his consent.

Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.radcr.2022.05.028.

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