Carcinoid Tumor of the Extrahepatic Bile Duct in an Adolescent: Prolonged Disease-free Survival After Surgical Resection

Tumor carcinoide de la vía biliar extrahepática en adolescente: larga supervivencia libre de enfermedad tras resección quirúrgica

Cancerous tumors of the extrahepatic bile duct are uncommon (less than 2% of all neoplasms). About 85% are cholangiocarcinomas, while the remaining 15% are miscellaneous tumors, including carcinoid tumors which represent 0.2%–0.34% of all bile duct neoplasms.\textsuperscript{1-7} The first case of carcinoid tumor of the bile duct was described by Davies in 1959, and up until 2009 only 60 cases had been published.\textsuperscript{1-4,6,8-9} We present a new case of bile duct carcinoma and discuss the symptoms, diagnosis, treatment and prognosis of this rare neoplasm.

The patient is a 14-year-old female with a self-palpated epigastric mass and abdominal distension. On examination, a mass in the right epigastrum and hypochondrium was detected. Results from blood tests (hemogram, hepatic biochemistry, coagulation, viral serology [hepatitis A, B, C and D; EBV, CMV] and tumor markers [CEA, CA19-9 and αFP]) were normal.

Ultrasound and abdominal CT showed a solid, isoechoic, homogeneous lesion in the hepatic hilum measuring 10 cm in diameter (Fig. 1). The tumor was in contact with and displaced the hepatic hilum and compressed the common bile duct, producing mild dilatation of the right intrahepatic bile duct. There were another 2 hyperechoic lesions measuring 14 and 27 mm in segment v.

With the diagnosis of probable hepatoblastoma, surgery was performed, and a large mass was observed in the hilar region that infiltrated the right liver lobe. Intraoperative ultrasound detected several lesions in segments νb, ν, νvi and νvii. Intraoperative biopsy was reported to be either hepatoblastoma or neuroendocrine tumor. A right hepatectomy was performed with resection of subsegment νb, portal lymph node dissection and total resection of the extrahepatic bile duct with hepaticojunostomy (Fig. 2).

During the postoperative period, the patient presented a low-output biliary fistula (grade A) that resolved with conservative treatment, and she was discharged on the 10th day post-op. The definitive histology results were: differentiated neuroendocrine tumor of the bile duct (carcinoid) with a moderate mitotic rate, and presence of numerous vascular thrombi with no lymph node involvement. Immunohistochemistry was positive for chromogranin, synaptophysin and PGPS; negative for vimentin, alpha-fetoprotein, alpha-antitrypsin, CEA, beta-HCG, 5-100 and neurofilaments; and (+/-) for EMA and PanCK. Postoperative determinations for 5 HIAA in urine as well as serum chromogranin A and serotonin were normal. One hundred months after the surgical procedure, the patient remains disease-free.

Carcinoid tumors arise from argentaffin cells (probably endodermal) that originate in the embryonic neural crest (Kulchitsky cells) and migrate during embryonic development.\textsuperscript{1-6,10} Carcinoid tumors of the biliary system (gallbladder and biliary tree) represent only 0.2%–2% of these lesions.\textsuperscript{1,2,4-8,9} This extremely low frequency is due to the fact that the settling of these cells in the bile duct is exceptional. It has been postulated that chronic inflammatory processes of the bile duct could induce phenomena of metaplasia in these cells and cause carcinoid tumor of the bile duct.\textsuperscript{1,4,10}

Carcinoid tumors of the bile duct are more frequent in women (ratio 2:1), and the mean age of presentation is 47 (range 10–79).\textsuperscript{1-3,5-8} There have only been 5 cases reported in adolescents and children.\textsuperscript{5,10} These tumors are located in: the common bile duct (55%), hilar region (30%), cystic duct (11%),

\textsuperscript{Please cite this article as:} Ramia JM, Garrote D, Muffak K, Villegas T, Ferrón A. Tumor carcinoide de la vía biliar extrahepática en adolescente: larga supervivencia libre de enfermedad tras resección quirúrgica. Cir Esp. 2014;92:636-638.
and hepatic duct (3%). The most frequent symptom is jaundice (55%–70%), followed by abdominal pain. Although they may be active hormonally, it is exceptional for them to present symptoms derived from hormonal secretion, and serum 5 HIAA levels are usually normal. Carcinoid tumors of the bile duct have been described in association with von-Hippel Lindau and MEN 1.

Correct preoperative diagnosis of these tumors is very uncommon. Several diagnostic methods are used (ultrasound, abdominal CT, MR cholangiography and ERCP). Intraluminal polypoid lesions on cholangiography are suggestive of carcinoid tumor of the biliary duct.

When a carcinoid tumor is found, there are 2 possible scenarios: it is usually (75%) an incidental finding during laparotomies performed for a misdiagnosis (choledocholithiasis, cholangitis), in the remaining cases, the intervention is done under the suspicion of cholangiocarcinoma. Carcinoid tumors of the bile duct have a series of typical clinical characteristics, including: female sex, 5th decade of life, limited local aggressivity, low rate of metastatic disease (30% of patients), and simple R0 resection.

In the literature, all the patients with R0 resection are alive and disease-free. Among those who present metastatic disease, 50% die during follow-up; their 5-year disease-free survival ranges between 40% and 80%, with a mean survival of 90 months. Postoperative treatment with chemotherapy (5-fluorouracil and streptozotocin) has not been able to demonstrate an increase in survival and, according to the NCCN, it is not indicated in completely resected cases.

**References**

Malignant Rhabdoid Tumor of the Colon: A Case Report

Tumor rabdoide maligno de colon: a propósito de un caso

Malignant rhabdoid tumors are highly aggressive neoplasms that were initially described in the kidneys of children as a rare variation of Wilms tumors with a rhabdomyosarcomatoid pattern and particularly poor prognosis.\textsuperscript{1} Subsequently, tumors with histologically similar characteristics were found in other locations, and classified as extrarenal malignant rhabdoid tumors (EMRT).\textsuperscript{2,3} These neoplasms are rare and have a very aggressive behavior. Their origin has been debated, as they have been described in several solid organs. To date, only 43 cases have been reported in the gastrointestinal tract, 5 the esophagus, 16 the stomach, 10 the small bowel and 12 the colon.\textsuperscript{4}

We present the case of a 77-year-old male with a prior history of ischemic cardiopathy who reported having abdominal pain and rectal bleeding during the course of the previous month. Colonoscopy revealed a stenosing neoplasm with partial necrosis that was 78 cm from the anal margin. Biopsy demonstrated the presence of atypical cells compatible with carcinoma, although immunohistochemistry showed negativity of the tumor cells for CDx-2 and CK 20, which suggested a non-colonic origin. Pre-operative CT showed no evidence of distant disease (Fig. 1). During surgery, a large tumor was found in the descending colon that infiltrated the omentum and parietal peritoneum; a left hemicolecotomy with primary anastomosis and resection of the affected abdominal wall was performed. The pathology study determined the lesion was a high-grade infiltrating malignant neoplasm (pT4aN1bMx), with co-expression of cytokeratin cocktail and vimentin (Fig. 2A and B), with negative CK7 and CK20, CDx2 (−), ALC (−), desmin (−), BerEP4 (−), p53 (−), CD117 (−) and calretinin (−), compatible with primary rhabdoid tumor of the colon. On the 7th day post-op, the patient needed a reoperation due to anastomotic leak, which included resection of the anastomosis and terminal colostomy in the left lower quadrant. The patient’s postoperative recovery was slow and required a stay in the ICU. One month after the initial intervention, a follow-up CT reported several mediastinal and retroperitoneal lymphadenopathies. In addition, in the previous tumor bed, a focal hepatic lesion was found in segment VIII, suggestive of metastasis, along with ascites and radiological signs of

\begin{figure}
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\includegraphics[width=\textwidth]{figure1.png}
\caption{CT: Tumor in the descending colon with reaction of the perilesional mesenteric fat and in close relationship with the parietal peritoneum.}
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\section*{References}


\section*{Acknowledgments}

Presented as a poster at the 17th National Meeting of the Spanish Society of Coloproctology in Palma de Mallorca from May 8 to 10, 2013.