Solid pseudopapillary tumor of the pancreas (SPPT). Still an unsolved enigma

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ABSTRACT

Solid pseudo-papillary tumor (SPPT) is a rare cystic tumor of the pancreas (1-3% of exocrine tumors of the pancreas) which shows an “enigmatic” behavior on the clinical and molecular pattern. A retrospective analysis of the citological studies and resected specimens of pancreatic cystic tumors from May 1996 to February 2010 was carried out. Three cases of SPPT were found, which are the objective of this study. In the three cases the pre-operative diagnosis was confirmed by cytology and specific immunohistochemical staining. Cases 2 and 3 showed strong immunoreactivity for Beta-Catenin and E-Cadherin staining. Radical resection (R0) was carried out in the three cases. A young male –21 years of age (case 1)– who had duodenal infiltration and two lymph nodes metastases died of hepatic and peritoneal recurrence 20 months following surgery. The other two cases are free of disease. The current review of the literature reports roughly 800 cases since the first report in 1959, and shows the enigmatic character of this tumor regarding the cellular origin, molecular pathways, prognostic factors and clinical behavior.

Key words: Solid-pseudopapillary tumor. Pancreatic cystic neoplasm. Prognosis. Beta-catenin.

RESUMEN

El tumor pseudopapilar (TSPP) es un tumor quístico del páncreas muy poco frecuente (1-3% de los tumores exocrinos del páncreas) y que tiene un comportamiento oncológico y molecular “enigmático”. Se realizó un análisis prospectivo de las citologías de las lesiones quísticas del páncreas, así como de los tumores quísticos resecados entre mayo de 1996 y febrero de 2010, encontrándose tres tumores SSPP, motivo de este estudio. En los tres casos el diagnóstico fue ocasional en el TC abdominal a pesar de presentar unos tamaños entre 3 y 6 cm de diámetro. En los tres casos se confirmó el diagnóstico preoperatorio mediante citología e inmunohistoquímica. En los casos 2 y 3 se confirmó la positividad para Beta-Catenina y E-Cadherina. En todos los casos se realizó cirugía radical (R0). Un varón de 21 años –caso 1– que presentaba infiltración duodenal y metástasis ganglionares falleció de progresión hepática y peritoneal a los veinte meses de la cirugía. Las otras dos pacientes –casos 2 y 3– se encuentran libres de recidiva. La revisión de la literatura desde su descripción en 1959 hasta la actualidad –aproximadamente 800 casos– confirma el carácter enigmático de este tumor en relación a su origen celular, molecular, factores pronósticos y comportamiento clínico.

Palabras clave: Tumor sólido-pseudopapilar, Tumores quísticos de páncreas. Pronóstico.

INTRODUCTION

The solid pseudopapillary tumor (SPPT) of the pancreas was first described by Franz in 1959 and characterized by Hamoudi in 1970, resulted in either “Franz” or “Hamoudi” tumors (1,2). It is an uncommon pancreatic cystic tumor (1-3% of exocrine pancreas neoplasms) which shows an enigmatic pattern from oncological and molecular perspectives. Long-term survival superior to 10-12 years have been reported in spite of vascular invasion and peritoneal or hepatic involvement (3-5). The SPPT usually has a benign evolution affecting young females (2nd-3rd decade), having related with sex hormones etiology although this hypothesis has not been confirmed. Some authors have reported a more aggressive evolution in young males (6-8).
Its biological and clinical pattern differs from the rest of the cystic neoplasms of the pancreas, which have aroused a great interest in the last decade (9-11). Since 1933, 718 cases have been described until 2005 and currently around 800 cases have been reported (10-17).

We present three cases of SPPT with different clinical features and outcomes, which confirm the “enigmatic” characteristics of this tumor which has been occasion of an editorial in this journal in 2006 (9).

PATIENTS AND METHODS

Between October 1996 and February 2010, 118 cytologic studies of cystic lesions of the pancreas were performed in our Institution. Most of them were obtained by fine-needle aspiration (FNA) during computerized tomography (CT) or upper endoscopic ultrason sound (EUS) procedure. In October 2006 the brush cytology technique was introduced (Echobrus Cook Medical, Winston-Salem, North Carolina, USA). A retrospective analysis of the clinical, surgical and cytopathological parameters of the cystic tumors diagnosed according with the 1996 World Health Organization (WHO) guidelines and reviewed in 2000 was performed (18,19). Three cases of SPPT were diagnosed in that period.

Case 1. A 17 year-old male with previous history of duodenal ulcer, presented a great ulcer and deformation of duodenal wall in a routine endoscopy. An abdominal CT and magnetic resonance imaging (MRI) showed a great solid-cystic lesion in the head of the pancreas. The FNA cytology reported an “epithelial malignant tumor”. The patient was referred to our Center. On admission, the blood analysis showed: Hb. 9.3 g/dL; Hto. 27.5%; Iron: 11 µg/mL. Total Bilirubin: 4.54 mg/dL; Gastrin 101 pg/mL (normal values < 150 pg/mL) and CA-19.9: 11.3 U/mL.

In the abdominal CT a 6 cm diameter tumor in the head of the pancreas was assessed, with compression on the duodenal wall and impingement of the mesenteric vein and celiac axis as well. The FNA cytology was informed as “solid-pseudopapillary neoplasm” with pseudopapillary pattern.

The patient underwent a total pancreactectomy due to global involvement of the pancreatic gland by a solid-cystic tumor without infiltration of the mesenteric vessels. The patient was reoperated at the 11th postoperative day for a gastrojejunal dehiscence and was discharged on the 20th postoperative day. The gross description reported a solid mass of 6.5 cm diameter which involved whole gland with intraluminal invasion and multiple hemorrhagic cystic structures. The microscopic study described a neofomation with papillary pattern alternating with microcystic areas. The tumor invaded the duodenal wall with negative surgical margins. Two of the lymph nodes were infiltrated. The patient was treated with adjuvant radiotherapy (51 Gy) and chemotherapy (Cysplatin, 5-FU) and later on with Docetaxel and Gemcytabin (20). 14 months after operation a loco-regional recurrence and liver metastases were observed dying 20 months after surgical resection.

Case 2. An incidental 6 x 6.5 cm diameter cystic lesion located in the tail of the pancreas was discovered in asymptomatic 28 year-old female. The EUS confirmed the expansив pattern. The cytopathological study described several branch of papillary fronds, consisting of a central fibrovascular stalk covered with several layers of neoplastic cells compatible with SPPT. The patient underwent laparoscopic distal pancreatectomy without incidences and remains free of recurrence and symptoms 4 years after operation. The gross appearance revealed a well circumscribed solid-cystic lesion of 6.5 cm diameter, with multiple hemorrhagic and necrotic areas (Fig. 1-A). The microscopic study revealed an epithelioid neoplastic alternating solid pattern with cystic spaces and pseudopapillae formed by connective tissue stalks which delineated clusters of epitheloid cells. The surgical margins and lymph nodes were negative (Fig. 1-B).

The immunohistochemical study was positive for Vimentin, α1-antitrypsin (AAT), CD-10 and CD-56; and negative for cytokeratin-7, cytokeratin-19, synaptophysin and chromogranin A.

Case 3. A 24 year-old female who consults for a second opinion with the previous diagnosed of pancreatic cystoadenoma. She was diagnosed 8 years before with occasion of an incidental finding in an abdominal CT when the lesion measured 3 cm diameter size.

In posterior controls she remained asymptomatic although an increase of the lesion up to 4.5 cm diameter and portal vein compression was assessed when she decided to consult in our Department (Fig. 2).

The SPPT diagnosis was confirmed by EUS and FNA cytology. The immunohistochemical staining was positive for nuclear beta-catenin and cytoplasmatic CD-10 (Fig. 3). The patient underwent Whipple procedure and discharged on 5 postoperative day with regular diet. The gross study described a well circumscribed solid lesion of 3.5 x 3.5 x 4.5 cm size in the head of the pancreas which displaced the pancreatic parenchyma and obstructing the Wirsung duct (Fig. 4). In the microscopic study, an epithelioid neoplasia growing in cellular clusters delineated by connective tissue stalks with pseudopapillary structures.

The tumor was well circumscribed by a fibrous tissue band with surgical margins free of disease and negative nodes. The immunohistochemical study was positive for CD-10, nuclear and cytoplasmatic Beta-Catenin and for progesterone receptors (85%).

DISCUSSION

The SPPT was first described by Franz in 1959 and posteriorly characterized by Hamoudi in 1970 (1, 2).
The tumor was denominated solid-pseudopapillary with occasion of the exocrine pancreas classification in 1996 (18). It is a rare tumor (1-3% of the exocrine neoplasms) which predominantly affects young females between 20-30 years of age, and with an overall 5 years survival rate of 95-98% and 10 year of 93% respectively after surgery (12,22).

From the biological, molecular and oncological point of view, it represents an “enigmatic” tumor from which the origin, prognostic factors and natural history are unknown, having aroused a great interest and awareness in the last decades, when several clinical series and excellent reviews have been reported (4,8-11,13,15-17,22-26).

From the three presented cases, highlights the first one, the case of a 17 year-old boy whenever the prevalence in males is between 3.9% and 6.6% (12,22). Machado and cols. reported the tumor in 7 males in a case series of 34 patients. This author reported a more aggressive histological pattern in the males although not definitive prognostic factors were obtained due to the limited number of patients (8). Other authors have reported bigger and more solid tumors in males without differences in survival (6,7). Table I summarizes the anatomical and clinical features of the largest series.

In the previous mentioned patient, the clinical presentation and evolution after radical surgery (R0) was very...
unusual developing early loco-regional recurrence and distant development of metastases in spite of adjuvant treatment, dying on the 16 postoperative month. We have found ten cases of a total of 433 published patients (2.3%) in 17 clinical series with a similar evolution (4,6;8,11,14,16,23-32). Tang et al., from Memorial Sloan-Kettering, published two similar cases out of 36 patient series, who died at 6 and 16 months after surgical resection. They were two females of superior age to the mean (33 and 45 years) who had large tumors of 9 and 20 cm diameter size respectively. One case showed a synchronic liver metastases and the other one lymph node involvement. Tumor necrosis, diffuse invasion, a higher mitotic index and infiltration with invasion of the duodenal lumen, as well as in our case number 1, were the differential findings (11,33) with the remaining 34 patients.

One of the paradigmatic fact of SPPT is that long-term survival—superior to 5 – 12 years—have been assessed in the setting of patients with liver metastases, multiple

Fig. 3. Fine-needle aspirate from a patient with SPPT (case 3). A. Demonstrates fibrovascular stalks with cells lined at the surface an periphery of the core (Romanowsky stain x 200). B. Demonstrates ovoid cells lining the fibrovascular core with finely granular chromatin pattern (Romanowsky stain x 400). C. Detail showing the nuclear pattern with intense labeling for β-Catenin (x 400). D. Demonstrates positive staining with immunoperoxidase for CD-10 (x 400) from the same case.

Fig. 4. Gross appearance of pancreatoduodenectomy specimen for SPPT (case 3) showing the well circumscribed solid lesion in the head of the pancreas, displacing the rest of the parenchyma.
node metastases and incomplete resections as well (10,12,14,16).

In our cases, the diagnosis was established as incident al lesion on CT imaging in spite of being large tumors with an axial diameter between 4.5 and 8 cm. When presenting, the abdominal symptoms are vague and unspecific (epigastric discomfort, nausea, weight loss). Only 1% of the patients present jaundice, even with tumors located in the head of the pancreas, as occurred in cases 1 and 3 (10,12,16,17,22).

The radiological features on CT and MRI show well circumscribed lesions with solid cystic changes and the presence of hemorrhagic areas within the tumor with high signal intensity on T1-weighted images after gadolinium –enhanced pulse sequences- (34). The three cases were preoperatively diagnosed with the FNA cytology study. This method, which is the most accurate for the cystic neoplasms of the pancreas, has been unusually described by most of the authors. In a review of 718 patients, only 52 cases (7%) were diagnosed by the FNA (12). Salvia et al. performed this technique in only 10 patients out of 31 (32.2%) SPPT’s patients with a sensitivity of 50% (14).

Jani reports a 75% efficacy (21/28 cases) with FNA guided in EUS exploration and points out the value of the immunohistochemical study in the cytology (35,36).

From the cytological view the different diagnosis with neuroendocrine tumors (NET) it could be difficult, whereas a positive staining for chromogranine A, CD10, PR and CD56 have been described in both tumors (11,37-39).

The nuclear and cytoplasmic expression of E-Cadherin and beta-catenin members of the Wnt pathway as specific markers have been reported in the 100% of TSP meanwhile on the NET the beta-catenin expression is located in the cellular membrane (24,40-42). These techniques should be incorporated in the diagnostic armamentarium when SPPT would be suspected (43). In our most recent case –number 3- the nuclear immunoreactivity for Beta-Catenin confirmed the diagnosis.

The origin of the SPPT is still an enigma. The original cell of the tumor has not been identified in spite of the recent immunohistochemical findings (44). Some authors have proposed the ovarian origin due to the relation of the pancreas with the genital ridge during the early embryogenesis (44). Geers has recently suggested the pancreatic acinar cells as its origin, although the debate regarding the SPPT ontogenesis is continuing (9,40,41,45,46).

The prognostic factor neither have been identified, having related the outcomes with the tumor size, mitotic index, necrosis grade, duodenal invasion and the presence of lymphatic metastases (8,11,16,47). This enigmatic pattern contrasts with the recent knowledge of the cystic tumor of the pancreas as the intraductal papillary mucinous neoplasm (IPMN) (48,49,50).

Finally we mention the alteration on β-Catenin / E-Cadherin. The β-Catenin protein is involved in the development and organogenesis of multiple tissues regulated by the Wnt/β-Catenin signalization pathway. Recent studies have pointed out the role of this pathway in the pancreas organogenesis being inhibited in the adult pancreas tissue (51,52). β-Catenin/APC mutations and Wnt signaling pathway alterations have been associated with the pancreaticoblastoma, acinar carcinoma, pancreatic ductal adenocarcinoma and SPPT (40,41,53-55).

Heiser et al. has recently developed an experimental model in mice, inducing the mutation in that pathway, suggestion that these alterations would be the origin of SPPT (46). In spite of these findings the SPPT is an “enigmatic” tumor with aspects remaining to be revealed.

### Table I. Clinical experience in SPPT*. Most numerous series in adults

<table>
<thead>
<tr>
<th>Author (Ref.)</th>
<th>Year</th>
<th>Age (Range)</th>
<th>No. cases</th>
<th>Sex ratio (M/F)</th>
<th>No. (%)</th>
<th>Symptoms (n (%))</th>
<th>Size (cm)</th>
<th>DPC</th>
<th>PD</th>
<th>PC</th>
<th>PT</th>
<th>Enuc.</th>
<th>Other</th>
<th>Follow-up (m)</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sun (23)</td>
<td>2003</td>
<td>27 (11-59)</td>
<td>28</td>
<td>23/5</td>
<td>23 (82%)</td>
<td>6 (1,2-6)</td>
<td>10</td>
<td>16</td>
<td>2</td>
<td></td>
<td></td>
<td>67 (7-110)</td>
<td>28 alives</td>
<td>1 liver recurrence</td>
<td></td>
</tr>
<tr>
<td>Papavramidis* (12)</td>
<td>2005</td>
<td>22 (2-89)</td>
<td>710</td>
<td>626/84</td>
<td>6.08 (0.5-34)</td>
<td>142</td>
<td>221</td>
<td>7</td>
<td>5</td>
<td>26</td>
<td>150 *</td>
<td>—</td>
<td>87 (3-240)</td>
<td>95% alive at 5 y.</td>
<td></td>
</tr>
<tr>
<td>Timptom (27)</td>
<td>2006</td>
<td>30 (15-57)</td>
<td>14</td>
<td>13/1</td>
<td>nc</td>
<td>7 (4-16)</td>
<td>3</td>
<td>10</td>
<td></td>
<td></td>
<td></td>
<td>87.6 (3-240)</td>
<td>34 alives</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>Salvia (14)</td>
<td>2007</td>
<td>31 (7-56)</td>
<td>31</td>
<td>27/4</td>
<td>14 (45%)</td>
<td>6 (2-15)</td>
<td>9</td>
<td>11</td>
<td>6</td>
<td>1</td>
<td>4</td>
<td>58.2 (7-229)</td>
<td>62 alives</td>
<td>2 rec. at 16 m. &amp; 7 y.</td>
<td></td>
</tr>
<tr>
<td>Lee (4)</td>
<td>2008</td>
<td>30 (18-63)</td>
<td>62</td>
<td>57/5</td>
<td>21 (33%)</td>
<td>6 (1.5-14)</td>
<td>19</td>
<td>38</td>
<td>2</td>
<td>1</td>
<td>2</td>
<td>47.5 (5-240)</td>
<td>34 alives</td>
<td>1 rec. at 7.7 y.</td>
<td></td>
</tr>
<tr>
<td>Machado (8)</td>
<td>2008</td>
<td>23 (10-72)</td>
<td>34</td>
<td>27/7</td>
<td>21 (79%)</td>
<td>7 (1.5-15)</td>
<td>11</td>
<td>15</td>
<td>5</td>
<td></td>
<td></td>
<td>57.6 (6-308)</td>
<td>34 alives</td>
<td>No recurrence</td>
<td></td>
</tr>
<tr>
<td>Reddy (16)</td>
<td>2009</td>
<td>32 (13-75)</td>
<td>37</td>
<td>33/4</td>
<td>27 (87%)</td>
<td>6.0 (0.3-12)</td>
<td>14</td>
<td>20</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>57.6 (0.8-328.8)</td>
<td>34 alives</td>
<td>No recurrence</td>
<td></td>
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*Review. DPC: Pancreaticoduodenectomy. DP: Distal pancreatectomy. CP: Central pancreatectomy. TP: Total pancreatectomy. Enuc.: Enucleation
REFERENCES


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