Spontaneous rupture of a non-functioning pancreatic neuroendocrine tumor
A case report of a rare cause of acute abdomen


*University of Milan, Fondazione IRCCS Ca’ Granda Ospedale Maggiore Policlinico, Milano (MI), Italy
** Department of General Surgery, Ospedale Manzoni, Lecco, ASST Lecco, Italy
***Department of Surgery, University of Milan-Bicocca, Istituti Clinici Zucchi, Monza, Italy

Spontaneous rupture of non-functioning pancreatic neuroendocrine tumor. A case report and review of the literature

BACKGROUND: Pancreatic neuroendocrine tumors (PNETs) are a group of rare tumors that account for 2% of all pancreatic malignancies, even though their incidence has been increasing over the past 20 years. Most PNETs are sporadic and tend to affect older individuals. Differently from functional tumors, which present with symptoms resulting from the specific hormone being elaborated, non-functioning pancreatic neuroendocrine tumors (NF-PNETs) typically present with symptoms related to local mass effect or metastatic disease. Today, due to the increasing use of abdominal imaging, NF-PNETs are frequently discovered incidentally.

CASE REPORT: A 32-year-old woman was admitted to our emergency department for worsening upper abdominal pain radiating to the back. Shortly after the admission, shock and peritoneal signs developed. An abdominal computerized tomography scan showed a solid mass (9 x 12 cm) of the pancreatic tail with severe hemoperitoneum. Exploratory laparotomy and subsequent distal splenopancreatectomy were performed for a bleeding tumor. Histopathological report showed a neuroendocrine, well differentiated tumor (G1). The postoperative course was uneventful and the patient was successfully discharged on 10th postoperative day.

CONCLUSION: Spontaneous rupture of solid neuroendocrine neoplasms of the pancreas can cause acute abdomen with potentially devastating effects.

KEY WORDS: Abdominal pain, acute abdomen, Hemorrhagic shock, Pancreatic neuroendocrine tumor

Background

Pancreatic neuroendocrine tumors (PNETs) comprise a heterogeneous group of neoplasms that arise from pluripotent cells in the pancreatic ductal/acinar system with variable clinical expression and progression. The PNET incidence is approximately 0.43/100,000 per year, accounting for 7% of all NETs, and has been increased in the last two decades. This growing incidence of PNET can be explained by the increased frequency of abdominal imaging, specifically computed tomography (CT), which leads to a high rate of asymptomatic tumors incidentally diagnosed. Clinically, PNETs are often classified as functional or non-functional based on the presence or absence of a particular clinical syndrome associated with hormone hypersecretion. Differently from functioning neoplasms, non-functioning pancreatic neuroendocrine tumors (NF-PNETs) do not cause paraneoplastic symptoms and usually present with symptoms of mass effects, distant metastases, or both. Consequently, NF-PNETs are often latey diagnosed due to their vague symptoms and rarity. Although surgical resection is considered the mainstay of curative treat-
ment for NF-PNETs, for small tumors less than 2 cm with a low proliferative index (Ki-67) observation may be an appropriate option. The overall 5-year survival rate is approximately 30%, with grading and systemic metastases at presentation as most significant prognostic factors. Few cases of spontaneous ruptures of pancreatic tumor have been reported in the literature and none of them concerns a NF-PNET. Herein, we present the first case of spontaneous rupture of NF-PNET leading to acute abdominal pain and hemorrhagic shock.

Case Presentation

A 32-year-old woman was admitted to our Emergency Department with severe abdominal pain in the left hypocondrium radiating to the back that had begun two hours earlier. She had a history of iron deficiency anemia successfully treated and reported multiple episodes of diarrhea in the last two months. Physical examination revealed abdominal distention with tenderness in the left upper quadrant. Vital signs were the followings: pulse rate 100 beats/minute, blood pressure 115/70 mm/Hg and body temperature 36.5 °C. Blood tests showed decreased hemoglobin concentration (9.5 g/dl) with elevated levels of white blood cells (13,000 per ml), normal level of C-reactive protein, normal platelet count and creatinine level. Liver enzyme, alkaline phosphatase and serum amylase were within the normal limits. One hour after admission the patient’s general condition rapidly deteriorated with increasing abdominal pain in the upper quadrants and pre-syncopal symptoms. Blood pressure decreased to 90/60 mm/Hg, heart rate increased to 130 beats/minute and arterial blood gas analysis showed a further decrease of hemoglobin level (8.1 g/dl). Urgent abdominal CT scan showed a solid mass (9x12x8 cm) with some cystic components in the pancreatic tail, associated with a massive hemoperitoneum and splenic vein compression. Also ecstatic collateral venous vessels along the course of the gastroepiploic vein and the mesenteric vein were reported (Fig. 1). Considering the rapid development of anemia and the deterioration of clinical conditions, the patient underwent emergency surgery. An exploratory laparotomy was performed, showing the presence of blood clots in the abdominal cavity and active bleeding from a ruptured mass in the pancreatic tail. A distal splenopancreatectomy was necessary, since the spleen was displaced cranially and the hilar region was infiltrated by the tumor (Fig. 2). In the following days only one blood transfusion was needed and the concentration of the amylase in the fluid from the peritoneal drainage rapidly normalized. Total length of hospital stay was ten days. Definitive histopathologi-
cal examination revealed a neuroendocrine, well-differentiated tumor (G1) with a mitotic count <2 per 10 high power fields (HPF), a Ki67 index lower than 3% and free surgical margins (Fig. 3). During surgical follow-up the patient underwent a scintigraphy with a somatostatin analogue (OctreoScan®) which did not show evident pathological uptake of the tracer in the supra and sub-diaphragmatic sites. Fourteen months postoperatively, there was no evidence of recurrence.

Discussion and Conclusions

NF-PNETs account for more than 75% of all pancreatic neuroendocrine tumors and represent a heterogeneous group of neoplasms with uncertain behavior. Nowadays, due to the widespread use of abdominal imaging, most of NF-PNETs are accidentally discovered, with more than 60% presenting with synchronous liver metastases. Less often patients present with nonspecific symptoms related to tumor invasion or displacement of contiguous structures, including jaundice, abdominal pain, weight loss, abdominal mass, nausea and vomiting. Acute abdomen is not considered as a possible clinical manifestation of this type of tumor, indeed no case of spontaneous bleeding or rupture is reported in the literature. Spontaneous abdominal hemorrhage is defined as non-traumatic and non-iatrogenic intra-abdominal bleeding and can be caused by the rupture of visceral organs. The sources of bleeding are often hypervascular neoplasms, such as hepatic adenoma or hepatocellular carcinoma, renal cell carcinoma or angiomylipoma and adrenal pheochromocytoma. Since the spontaneous rupture of a pancreatic neoplasm is an extremely rare event, we research the English literature for previous reported cases. We considered only cases of spontaneous ruptured of primary pancreatic neoplasm confirmed histologically. Ruptures occurred after traumatic events or cases of pancreatic metastases from other primary malignancies were excluded. A total of 13 cases of spontaneous ruptures of primary pancreatic tumors were identified. Patients and tumor characteristics are summarized in Table I. Tumor location in the pancreas was variable with most tumors found in the body or tail of the pancreas and only two cases found in the head. Mean tumor size was 11 cm, with all but one tumor being smaller than 8 cm in diameter. Regarding the tumor type, there was a predominance of two differ-

Table I - Reported cases of spontaneous rupture of primary pancreatic neoplasm.

<table>
<thead>
<tr>
<th>Canali EPase Number</th>
<th>Authors</th>
<th>Year of publication</th>
<th>Gender</th>
<th>Age</th>
<th>Tumour size (cm)</th>
<th>Type of the tumor</th>
<th>Location of the tumor</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Bombi JA et al.</td>
<td>1984</td>
<td>F</td>
<td>22</td>
<td>12</td>
<td>Solid pseudopapillary neoplasm</td>
<td>body</td>
</tr>
<tr>
<td>2</td>
<td>Annali Ep &amp; Achilli al.</td>
<td>1986</td>
<td>F</td>
<td>33</td>
<td>10</td>
<td>MCN with associated invasive carcinoma</td>
<td>tail</td>
</tr>
<tr>
<td>3</td>
<td>Todani T et al.</td>
<td>1988</td>
<td>F</td>
<td>16</td>
<td>8</td>
<td>Solid pseudopapillary neoplasm</td>
<td>tail</td>
</tr>
<tr>
<td>4</td>
<td>Panieri E et al.</td>
<td>1998</td>
<td>F</td>
<td>34</td>
<td>12</td>
<td>Solid pseudopapillary neoplasm</td>
<td>body</td>
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<tr>
<td>5</td>
<td>Ozden S et al.</td>
<td>2007</td>
<td>F</td>
<td>32</td>
<td>15</td>
<td>MCN with associated invasive carcinoma</td>
<td>tail</td>
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<tr>
<td>6</td>
<td>Bergenfeldt M et al.</td>
<td>2009</td>
<td>F</td>
<td>42</td>
<td>11</td>
<td>MCN with associated invasive carcinoma</td>
<td>body</td>
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<tr>
<td>7</td>
<td>Naganuma S et al.</td>
<td>2011</td>
<td>M</td>
<td>39</td>
<td>12</td>
<td>Acinar cell carcinoma</td>
<td>head</td>
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<td>8</td>
<td>Mohammad A et al.</td>
<td>2012</td>
<td>M</td>
<td>5</td>
<td>12</td>
<td>Pancreatoblastoma</td>
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<td>9</td>
<td>Honda S et al.</td>
<td>2013</td>
<td>M</td>
<td>13</td>
<td>4</td>
<td>Solid pseudopapillary neoplasm</td>
<td>tail</td>
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<tr>
<td>10</td>
<td>Takamatsu S et al.</td>
<td>2014</td>
<td>F</td>
<td>12</td>
<td>13</td>
<td>Solid pseudopapillary neoplasm</td>
<td>body</td>
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<tr>
<td>11</td>
<td>Pattanshetti VM et al.</td>
<td>2015</td>
<td>F</td>
<td>31</td>
<td>-</td>
<td>Solid pseudopapillary neoplasm</td>
<td>body</td>
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<tr>
<td>12</td>
<td>Omori H et al.</td>
<td>2015</td>
<td>M</td>
<td>69</td>
<td>-</td>
<td>Ductal adenocarcinoma</td>
<td>body</td>
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<tr>
<td>13</td>
<td>Narita M et al.</td>
<td>2015</td>
<td>F</td>
<td>32</td>
<td>12</td>
<td>Neuroendocrine tumor</td>
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<td>14</td>
<td>Our patient</td>
<td>2019</td>
<td>F</td>
<td>32</td>
<td>12</td>
<td>Neuroendocrine tumor</td>
<td>tail</td>
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</tbody>
</table>
ent histotypes: solid pseudopapillary neoplasm (SPN) and mucinous cystic neoplasm (MCN) with associated invasive carcinoma. MCNs of the pancreas often grow slowly and can remain indolent for many years \(^1\). Since the stroma usually expresses estrogen and progesterone receptors \(^2\)-\(^4\), MCNs that develop during pregnancy tend to be huge with a high risk of rupture and bleeding, as reported in three out of the four cases \(^10\),\(^13\),\(^15\). Thus, the possibility of ruptured pancreatic mucinous cystic neoplasms should be kept in mind, especially facing an acute abdomen in pregnancy. SPNs are low grade malignant tumor accounting for \(1-2\%\) of exocrine pancreatic neoplasms and typically present in young females as a large asymptomatic epigastric mass \(^5\). Clinical presentation as acute abdomen is considered rare for this tumor and the incidence of rupture in a large series was reported in \(2.7\%\) of \(292\) cases \(^6\). We found six cases of spontaneous ruptures of SPN reported in the literature, all of them occurred in young women with a mean age of \(21\) years \(^9\),\(^11\),\(^12\),\(^17\),\(^19\). The other histotypes reported in our review include ductal adenocarcinoma, acinar cell carcinoma and pancreatoblastoma \(^7\),\(^16\),\(^19\). To our best knowledge, our patient is the first case of spontaneous rupture of PNET reported in the literature. Regarding surgical treatment of the reported cases, distal pancreatectomy with splenectomy was performed in five cases \(^10\),\(^11\),\(^14\),\(^19\),\(^20\), while spleen preserving pancreatectomy was possible in four patients \(^12\),\(^13\),\(^16\),\(^18\). Whipple’s procedure (pancreatice-duodenectomy) was necessary in one case \(^15\); in the last three patients tumor enucleations were performed \(^7\),\(^8\),\(^17\). One patient underwent laparoscopic surgery, while the others were submitted to urgent laparotomies \(^17\). Laparoscopic technique has been increasingly applied in recent years in the treatment of body and tail pancreatic lesions, leading to a significantly reduction in morbidity with uncompromised oncological outcomes \(^21\). Even though this minimally invasive approach can be considered safe and adequate in elective circumstances, its role as emergency procedure should be further explored \(^24\),\(^25\).

Thus, in case of ruptured lesions with active bleeding, an open approach with emergent laparotomy is considered the best choice. It is noted that, when performing distal pancreatectomy, especially in young patients, the spleen should be preserved in order to avoid the asplenic complications that may add much morbidity as well as mortality. In our case, since the hilar region of the spleen was infiltrated by the tumor, a splenopancreasectomy was necessary in order to achieve a radical removal and to reduce the risk of local recurrence. In conclusion, we reported the extremely rare case of a spontaneous rupture of a NF-PNET as cause of hemorrhagic shock. Even if considered slow growing neoplasms usually late diagnosed, NF-PNET’s can be rare source of spontaneous intraab TABLE Idominal bleeding requiring emergency surgery.

**References**


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