Neuroendocrine Tumours of The Ampulla Of Vater

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Abstract

Background

Neuroendocrine tumors rarely occur in ampulla of Vater and their incidence in current increase, in particular malignant forms. The final diagnosis is done on histology, and sometimes it can be difficult to diagnose them preoperatively, as they present a clinical picture similar to the adenocarcinomas of this region.

Case presentation

Case 1
A 43-year-old woman with epigastric pain, serum alkaline phosphatase and γ-glutamine transferase high, endoscopic retrograde cholangiopancreatography (ERCP) showed choledochal metastasis on suspicous ampullary stenosis. Surgical exploration has demonstrated the presence of a mass of 4 cm at the expense of the ampulla of Vater. An ampullectomy was carried out, histological and immunological aspect was in favor of a well differentiated neuroendocrine carcinoma with expression of synaptophysin and chromogranin with a Ki 67 at 2%. The patient remained under good control with complete cure and without disease for 51 months.

Case 2
A 28-year-old female consultant for cholestatic jaundice associated with general asthenia, abdominal computed tomography-scanner (CT scan) has objectified a tissue lesion process of the vater bulb protruding into the duodenal lumen, a pancreaticoduodenectomy was performed, pathologic examination and immunohistochemistry of the surgical specimen was in favor of a well differentiated neuroendocrine tumor with expression of synaptophysin, chromogranin A and CKAEl/AE3 with a Ki 67 at 2%. The patient remained under good control with complete recovery and no signs of illness 19 months after the surgical procedure.

Conclusions : Ampullary neuroendocrine tumors (ANETs) are distinct entities that present clinically with jaundice. They positively stain chromogranin A and synapthophysin immunohistochemistry. The Pancreaticoduodenal resection appears to be the procedure of choice for most patients with ANET because this procedure removes all the potential tumor-bearing tissue.

Keywords

Neuroendocrine carcinoma - Ampulla of Vater - pancreaticoduodenectomy – prognosis.

I. Background

Neuroendocrine tumors of the ampulla of Vater are very rare, (Less than 100 published cases) [1, 2, 3]. In 2000, the World Health Organization revised the histopathological classification system for gastroenteropancreatic neuroendocrine tumors[4, 5]. The tumor neuroendocrine replaced the carcinoid term, which invokes a broad spectrum of appearance and of behavior histological and it is subdivided in two categories: benign, potentially malignant and malignant. They have a better prognosis than adenocarcinomas and good results have been reported for pancreaticoduodenectomy and local resection[2]. We report two new cases with ampullary neuroendocrine tumors and through the data of the literature we take stock of the various aspects of this rare entity.

II. Case presentation

Case 1
A 43-year-old woman followed for goiter, who presented 1 year before her admission chronic epigastric pain. Hepatic function tests showed marked elevations of alkaline phosphatase and γ-glutamine transferase. ERCP showed choledochal metastasis on a suspicious ampullary stenosis. Surgical exploration has demonstrated the presence of a mass of 4 cm at the expense of the ampulla of Vater. An ampullectomy was carried out, histological and immunological aspect was in favor of a well differentiated neuroendocrine carcinoma with expression of synaptophysin and chromogranin with a Ki 67 at 2%. The patient remained under good control with complete cure and without disease for 51 months.
transferase. The bili-Magnetic resonance imaging (bili-MRI) has objectified a choledochal enhancement. Endoscopic retrograde cholangiopancreatography (ERCP) showed choledochal metastasis on suspected ampullary stenosis. The duodenoscopy showed a polyploid rounded ampullar ulcer formation measuring 4 cm. An ampullectomy was carried out, the anatomicopathological and immunohistochemical examination returned in favor of a well differentiated neuroendocrine carcinoma of 2.3 cm with expression of synaptophysin, chromogranin and CKAE1/AE3 with a Ki 67 at 2%, with tumor free margin. No adjuvant treatment was indicated. A thoraco-abdominopelvic CT scan was produced two months later objectived an aerobilie with persistence of the 10 mm dilatation of the main bile duct. The patient presented a complete recovery and disease-free recovery 51 months after the surgery.

**Case 2**

A 28-year-old woman presented 4 months before admission with a generalized asthenia associated with cholestatic jaundice, abdominal CT showed a tissue lesional process of the vater bulb protruding into the duodenal lumen with thickening of the inner wall of the D1, it measures 35x27 mm (Fig. 1). Histopathologic examination of the biopsy in favor of an interstitial duodenitis with calcification. A pancreaticoduodenectomy was performed (Fig. 3), the anatomicopathological and immunohistochemical examination of the surgical specimen was in favor of a well differentiated neuroendocrine tumor of 2.7 cm with expression of synaptophysin, chromogranin A and CKAE1/AE3 with a Ki 67 at 3% (Figs 4 and 5). one of the twenty examined ganglions was involved. An abdominal computed tomography of control was performed 3 months later, which showed an infiltration of the operating area without mass or visible pathological contrast enhancement (Fig. 2). She remained disease free19 months after surgery.

*Figure 1:* Scannographic axial sectionshowing a tissue processcentred on the bulb of Vater enhancedaftercontrast, bending in the duodenal light withthikening of the internalwall of D1, itmeasures 35x27 mm

*Figure 2:* Scannographic axial sectionshowed an infiltration of the operating area without mass or visible pathological contrast enhancement

*Figure 3:* Macroscopic appearance of the CDP patch with a beige nodular ampullary tumor (forceps).
III. Discussion

Neuroendocrine tumours of the ampulla of Vater are very rare gastrointestinal malignancy, they are less than 2% of tumoursampullary and less than 1% of the gastrointestinal tumoursneuroendocrines. The World Health Organization (WHO) published in 2000 the latest classification which advocates distinguishing 3 categories: poorly differentiated endocrine carcinomas called small cell carcinomas, well-differentiated endocrine carcinomas called malignant carcinoids and well-differentiated endocrine tumors which are carcinoids [4]. The first two categories are considered to be malignant neoplasms, while well-differentiated endocrine tumors may have malignant or benign clinical signs. Our patients had well-differentiated endocrine carcinoma.

There is another method of classifying neuroendocrine tumors (NET) which is based on two criteria which are the presence of necrosis and mitotic activity. [6, 7] The well differentiated NETs, low-grade (G1) which reveal a lack of necrotic areas and less than 2 mitotic figures per 10 HPF. Neuroendocrine carcinoma moderately differentiated, intermediate grade (G2) that show the presence of necrotic foci or 2 to 20 mitotic figures per 10 HPF. While, the poorly differentiated neuroendocrine carcinomas, high Grade (G3) show the presence of necrotic foci and more than 20 mitotic figures per 10 HPF and [7].

The median age is 48.6 years and the sex ratio of women and men is 2.8: 1 based on data from 105 published cases in the literature. [8, 9, 10, 11, 12, 13, 14] The main symptom on the day of the consultation is Jaundice (60%), followed by pain (40%) and weight loss (10%). One of our patients presented with jaundice. Other signs may be observed such as acute pancreatitis in 3 to 6%, upper gastrointestinal bleeding in 3% [15, 16, 17]. The carcinoid syndrome is rare in patients with carcinoid of the ampulla of Vater without liver metastasis and this can be explained by made that the majority of the hormones produced by functional tumors are eliminated by the metabolism of the first passage in the portal system [18]. In 25.9% carcinoids of the ampulla of Vater are often associated with the von Recklinghausen disease [1,19]. Contrary to our 2 cases, the disease of von Recklinghausen was not associated.

Patients followed for carcinoid tumors of ampulla of Vater are different from those with carcinoid tumors of the jejunum and the ileum because they are in the majority of cases devoid of clinical and biological signs of the carcinoid syndrome. Symptoms of carcinoid syndrome such as flushing, asthma and diarrhea are described in only 2.8% (2 out of 71) of patients followed for papillary carcinoids.[1]Our patients have no symptoms of carcinoid syndrome.

The preoperative diagnosis of carcinoid tumors is only correct in 10 out of 71 patients that is 14% according to the data of the literature.[1]The diagnosis of carcinoid tumors is based on ERCP coupled with deep biopsies. The detection of lymph node metastases and the depth of invasion is essentially done by endoscopic
ultrasound. Once the diagnosis of ANET has been established, the demonstration of metastases is done by computed tomography and octreotidescintigraphy[20,1]. It should be noted that on imaging there are no specific features that can distinguish ANET from adenocarcinoma. In our cases the diagnosis was made by the biopsy of the ampoule.

Histologically, ANETs are small, solitary, polyoid and are covered with flattened mucosa. Microscopically the tumors are arranged in nests, microglandular, trabecular and rarely insular pattern. The tumor cells are small, uniform with scanty granular cytoplasm. The nuclei are regular, normochromic with scanty mitosis. The signs in favor of malign behavior are vascular invasion, gross local invasion or metastasiss[6, 21]. Immunohistochemical stainingis the main diagnostic method for ANET, they stain positively with chromogranin A in 92% of cases and neuron-specific enolase, synaptophysin and cytokeratin in 100%. Somatostatin is found in 58 to 67% of cases. Serotonin is secreted in 17% of cases as well as CCK. Insulin will stain positive in 25% of tumors. However, 13% are negative for any hormonal staining[20].

One of the prognostic factors of peripancreatic adenocarcinoma is tumor size. Unlike ANETs in size does not predict the metastatic potential. In a study, on patients with 2 cm ampullary NET, showed that the diagnosis of lymphoid nodal disease was positive in more than half of them. Metastases were present in 46% of tumors of 2 cm, in 50% of tumors of 1-2 cm and in 66% of tumors of 1 cm [22, 23]. Other cases reported by Makhlof et al. [24] of two tumors measuring less than 2 cm demonstrate metastases, while a 5 cm tumor did not show any evidence of metastatic disease. The conclusion of these two reports is that ANET metastasize in approximately 50% of the time independently of the size of the tumor.

In general, non-angio-invasive tumors less than or equal to 1 cm with 2 or fewer mitoses per 10 high-power fields (HPF) are considered benign, whereas the others show a high risk of malignancy. [4] ANETs are not angio-invasive, they measure 1 cm and show up to two mitoses per 10 HPF. [25] Contrariwise, the spread of the neoplasm to the muscularis, with a high degree of mitotic activity (3 or more mitoses per 10 HPF) and a diameter of 2 cm are considered as risk factors for the occurrence of metastases [26, 27, 28].

The treatment of ANET is controversial, and there is no general rule regarding their optimal treatment, as they are very rare tumors with unpredictable biological behavior and prognosis [22]. As it has been shown that the relationship between tumor size and the presence of metastatic ganglion has not been clearly established and therefore pancreaticoduodenectomy remains the treatment of choice for ANETs regardless of tumor size outside of metastasis at a distance [17, 29, 26]. Pancreaticoduodenectomy has been done in more than 50% of patients, according to a review of 105 reported cases of literature. [30] As has been described in the most recent series in which pancreaticoduodenectomy has often been adopted [17, 31]. Radical excision was performed in one of our patients in this report, with a well differentiated NET and she remained free of illness for 19 months after surgery.

Reduction surgery has an important place in large tumors and should be indicated for patients with hypersecreting tumors of hormones, even in cases of local or distant metastasis, as it has shown an improvement in survival rates of up to 80%. At 5 years old [21], the main disadvantage of local ANET resection is that it is technically difficult and associated with significant morbidity [22, 32]. Therefore, it is recommendable to avoid the risk of exposing patients to the risk of missing the resection of their metastatic lymph nodes, thus missing their cure by a reliable oncological surgery. In general, local excision is limited for tumors <2 cm, highly differentiated and slow-growing, and this for patients with high surgical risk and unable to tolerate further surgery. [16, 17]

The serum chromogranin A level is a tumor marker that reflects the tumor burden of NETs, and is a tool for detecting tumoral relapse, disease response to treatment, and progression. [33, 22, 34]

The most important prognostic factors that determine the survival of ANET are grade and distant metastases. Other features are incriminated but less important in long-term survival such as lymph node involvement, tumor size and resection margins. [6, 35] High-grade tumors show a 10-year survival rate of 15%, while low-grade NETs have a 5-year survival rate of 80% and a 10-year survival rate of 71%. [15, 33] Overall survival of the resected ampullary NETs is 81% according to the grade of the tumor, whose overall survival at 5 years of well-differentiated tumors is 90% and which decrease markedly for low-grade tumors [6, 15, 35, 33].

NET of the ampulla of Vater are distinct entities that present clinically with jaundice. They positive stain with chromogranin A and synaptophysininmunohistochemistry. ANETs without invasion and a diameter of less than 2 cm are considered a better prognosis tumor. Pancreatoduodenectomy should be the standard treatment for ANETs because of the poor accuracy of preoperative and intraoperative assessment of lymph node involvement and the high incidence of lymph nodal metastasis, even in small tumor smaller than 2 cm.
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Abbreviations
ERCP: Endoscopic retrograde cholangiopancreatography.
CT scan: Computer tomography scanner.
ANET: Ampullary neuroendocrine tumor.
WHO: World Health Organization
Bili-MRI: Bili-Magnetic resonance imaging
NET: Neuroendocrine tumor

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The authors declare that they have no competing interests.
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