Case Report

Tracking Down Duodenopancreatic Malignancy

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Background Malignant tumours of the duodenum are rare and often difficult to diagnose. Due to the small clinical experience with duodenal malignancies their prognosis is unknown and resection is the treatment of choice.

Case report Adding to a small series of incidental tumours, we report the case of a 65-year-old patient with primary extranodal (MALT-) lymphoma of the duodenum infiltrating the pancreatic head. The patient was admitted because of anaemia and epigastric discomfort with a history of Helicobacter-pylori associated gastric ulceration. Physical examination and bloodchemical values were otherwise normal. Endoscopy revealed duodenal ulceration but the biopsies taken from the ulceration did not give any evidence of malignancy or residual Helicobacter pylori infection. But MRT showed a circular intramural tumour of the duodenum. On laparotomy a large duodenal tumour adherent to the pancreatic head was found and a Whipple procedure was performed.

Conclusion Apart from describing the case of a rare lymphoproliferative disorder of the duodenum, this report illustrates the diagnostic difficulties with uncommon neoplasm's of the duodenopancreatic region and the value of MRT prior to resection of a duodenopancreatic mass.

Keywords: MALT-Lymphoma, duodenum, pancreas, MRT

INTRODUCTION

Malignant neoplasm's of the small bowel are rare and their prognosis is vague [1]. A recent epidemiological, US-American study [2] found an average annual incidence rate of 9.9 per million people for malignant tumours of the small intestines. The incidence was higher among men than among women and was found to be overall rising and rising with age. The most common histologic subtypes in this study were carcinoid tumours (3.8/million) and adenocarcinoma (3.7/million), sarcoma (1.3/million) and lymphoma (1.1/million). Smaller series have seen adenocarcinoma [3] or lymphoma [4] ahead in incidence. Adenocarcinoma seems to be more common in industrialised countries, where intestinal lymphomas are rather rare [5]. Intestinal lymphomas form a heterogeneous subgroup of malignant lymphomas currently classified in the Kiel-, or most recently in the Revised European - American Classification of Lymphoid Neoplasm's (=REAL) [6]. Intestinal B-Cell
lymphoma, most often large cell lymphoma [7], is common in the ileum [5] but rare in the duodenum and may mimic adenocarcinoma of the pancreatic head or duodenum or other neoplasms of the duodenopancreatic region [8].

Diagnosis of tumours of the duodenopancreatic region, as it has been experienced by various authors, is often obscured by vague abdominal symptoms e.g., abdominal discomfort, nausea, vomiting or weight loss. Physical examination may reveal a palpable mass in patients with advanced tumour growth.

Endoscopic diagnosis of duodenal lesions often fails because of a cryptic tumour site or an essentially submucosal or intramural tumour growth as in the presented case [9-11]. Brush cytology may be of value in tumours of the papilla of Vater but often is not conclusive when taken from other sites [12]. Biopsy specimens, too, fail to verify a histologic diagnosis due to endoscopically not accessible tumour sites. Laboratory examinations and contrast enhanced radiographs often lack characteristic findings as well.

Modern techniques of all-in-one MRT, MRCP and MRA may help to specify the diagnosis of tumours of the duodenopancreatic region [13] and to determine resectability [14]. Sensitivity of MRT is high but specificity varies with tumour site.

The case history, presented below, illustrates some of these diagnostic difficulties of characterising malignant tumours of the duodenopancreatic region and emphasises the value of high-quality MRT.

CASE REPORT

A 65-year-old man was admitted because of anaemia, loss of weight, fatigue and epigastric discomfort with a suspected tumour of the duodenopancreatic region. CT of the abdomen, obtained prior to admission, had shown a mass in the duodenopancreatic region, suspicious for pancreatic adenocarcinoma.

On admission the patient was pale with a haemoglobin of 8.4 g/dl. Bloodchemical values, including serum electrophoresis, immune electrophoresis, differential blood count, and physical examination were normal.

The patient had a history of gastric ulceration and had received eradication for Helicobacter pylori infection one year prior to admission. He was under Phenobarbital medication for post-traumatic epilepsy but had not had seizures for 10 years. He denied recent fever, loss of weight, nausea or vomiting, jaundice, changes in bowel habits or quality of stool and any other major diseases. He had been operated for bilateral inguinal hernia but did not have other abdominal surgery.

On gastroduodenal endoscopy the gastroduodenal passage appeared to be normal. An ulcer was found in the descending duodenum proximal of the papilla. Mucosal biopsies from this site showed signs of chronic inflammatory ulceration without Helicobacter pylori infection.

An endoscopic retrograde choledochopancreatography was performed demonstrating a regular shape of bile duct and main pancreatic duct.

Contrary to the endoscopic findings, hypotonic duodenography demonstrated subtotal stenosis of the descending duodenum (Fig. 1).

MRT and MRCP were performed to further characterise the suspected tumour. MRT (Fig. 2) demonstrated a segmental circular wall thickening limited to the descending duodenum. A submucosal tumour, e.g., lymphoma seemed to be the most likely diagnosis. The pancreatic head was normal in size. MRCP showed an unremarkable main pancreatic duct, excluding a pancreatic mass (Fig. 3).

Endoscopy was repeated and biopsies were taken from around the papilla of Vater. Histopathology of these biopsies again did not reveal any conclusive malignant findings.

On laparotomy a large duodenal tumour of circular intramural growth was found without
FIGURE 1  Hypotonic duodeno-radiography demonstrates a segmental stenosis of the descending duodenum.

FIGURE 2  Longitudinal MRT scan of the upper abdomen showing a segmental circular wall thickening limited to the descending duodenum. The pancreatic head is normal in size.
apparent infiltration of mucosa or serosa. The tumour, however, was found to infiltrate the papilla of Vater and the pancreatic head. Suspecting malignancy a Whipple procedure was performed. The macroscopic inspection of the specimen revealed a semicircular infiltrative mass in the duodenal wall (Fig. 4). The patient recovered well and was discharged 10 days after surgery. Adjuvant radiotherapy was performed starting four weeks after surgery.

Microscopic findings The tumorous mass measured 5 cm in longitudinal length and was located in the wall of the duodenum. The duodenal mucosa overlying the mass showed flat ulceration. Microscopically there was a transmural infiltration of the entire thickness of the duodenal wall by proliferating lymphoid cell. These cells infiltrated the serosa and the pancreatic parenchyma also (Fig. 5a). The papilla of Vater was free of blastomatous infiltrates.

The diffuse lymphoid infiltrates contained a large population of CD-20 positive cells. CD-3 positive small T-cells were present in smaller numbers and were dispersed in the B-cell proliferates. Ki67-index was low and κ-lightchains were restricted (Fig. 5b) in favour of λ-lightchains (Fig. 5c). These findings are conclusive of a low grade B-cell lymphoma according to the Kiel-classification scheme. The infiltration of lymphoid cells with lymphofollicular hyperplasia in the duodenal mucosa next to the mucosal ulceration, which merges with the blastomatous infiltration of low grade B-cell lymphoma in the underlying deeper zones of the duodenal wall, is highly suggestive of a B-cell lymphoma derived from mucosa-associated lymphoid tissue (=MALT) of the duodenum. Helicobacter pylori, however, was not evident neither in the tumour specimen nor in the gastroduodenal biopsies taken preoperatively.
FIGURE 4  Gross photograph of the duodenum with a segmental intramural tumour of circular growth. The papilla of Vater is indicated by a probe.

DISCUSSION

Reconsidering this case history, several questions arise: Could preoperative diagnosis been more effective, has surgery been necessary for this type of neoplastic lesion of duodenum and pancreas, and do primary (extranodal) duodenopancreatic lymphomas belong to the group of gastric lymphomas rather than to the group of intestinal lymphomas?

The few documented cases of primary (=extranodal) duodenal lymphoma reflect the variety of histologic subtypes of B-cell-lymphoma [15–17]. Lymphomas of the papilla of Vater may form another subgroup of intestinal B-cell-lymphoma [18–20]. Lymphoma of the stomach is known to occur in the setting of chronic Helicobacter pylori gastritis. Gastric lymphomas, located in the distal stomach, may invade the duodenum [21] and may mimic primary duodenal tumours there. In the presented case, duodenal ulceration and a history of Helicobacter pylori induced gastric ulceration may have been associated with the
pathogenesis of duodenal lymphoma. It has
to remain unclear, however, whether the re-
ported MALT-lymphoma represents a gastric
lesion, which had skipped into the duodenum
or a primary tumour.

Only little experience with the heterogeneous
group of small bowel B-cell lymphomas has
been gathered so far and treatment recommend-
dations are still a matter of debate [22]. Several
studies are currently on their way to compare
radiotherapy, chemotherapy, surgery and multi-
modal approaches. There is only minor evidence
[23], that eradication of Helicobacter pylori is
beneficial in other than gastric lymphoma. Re-
section still is the predominant treatment for
duodenopancreatic lymphoma [24, 25] as for any
duodenopancreatic tumour of suspected malig-
nancy, due to the lack of clinical data on medical
treatment options, and, of course due to the
difficulties of finding a preoperative diagnosis
as well. It is another "dilemma" for surgeons
[26] that the decision whether to resect or not
to resect such a tumour often has to be made
without a definite histopathologic diagnosis.

In the presented case, a tumour of the duo-
denopancreatic region was suspected but no
histologic findings were obtained in the course
of diagnostic procedures. Only MRT properly
identified an intramural duodenal tumour.
But experience in viewing such tumours still
is too limited as to provide a specific char-
acterisation of rare lesions. In such cases it

FIGURE 5 A photomicrograph of a low-power field of the tumour shows a diffuse proliferation of lymphoid cells with
infiltration of the pancreatic parenchyma (left field). Medium-power fields of this lymphoma, immunostained for κ-lightchains
(right upper field: negative) and for λ-lightchains (right lower field: positive in B-cells with lymphoplasmocytoid
differentiation) showing a lightchain restriction as an evidence for a neoplastic proliferation.
will be left to the surgeon to decide, whether to resect a duodenopancreatic tumour of unknown histology or not.

References


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