Case Report

Pancreatic non-functioning neuroendocrine tumor: a new entity genetically related to Lynch syndrome

Anna Serracant Barrera1, Sheila Serra Pla1, Carmen María Blázquez Maña2, Rubén Carrera Salas2, Neus García Monforte1, Natalia Bejarano González1, Andreu Romaguera Monzonis1, Francisco Javier Andreu Navarro2, María Rosa Bella Cueto1, Francisco G. Borobia1

1Department of Hepatobiliary Pancreatic Surgery, 2Department of Pathology, Parc Taulí, Hospital Universitari, Sabadell, Barcelona, Spain

Correspondence to: Francisco G. Borobia. Department of Hepatobiliary Pancreatic Surgery, Parc Taulí, Hospital Universitari, C/Taulí s/n, Sabadell, Barcelona 08208, Spain. Email: fgborobia@gmail.com.

Abstract: Some pancreatic neuroendocrine tumors (P-NETs) are associated with hereditary syndromes. An association between Lynch syndrome (LS) and P-NETs has been suggested, however it has not been confirmed to date. We describe the first case associating LS and P-NETs. Here we report a 65-year-old woman who in the past 20 years presented two colorectal carcinomas (CRC) endometrial carcinoma (EC), infiltrating ductal breast carcinoma, small intestine adenocarcinoma, two non-functioning P-NETs and sebomatricoma. With the exception of one P-NET, all these conditions were associated with LS, as confirmed by immunohistochemistry (IHC) and polymerase chain reaction (PCR). LS is caused by a mutation of a mismatch repair (MMR) gene which leads to a loss of expression of its protein. CRC is the most common tumor, followed by EC. Pancreatic tumors have also been associated with LS. Diagnosis of LS is based on clinical criteria (Amsterdam II and Bethesda) and genetic study (MMR gene mutation). The association between LS and our patient's tumors was confirmed by IHC (loss of expression of proteins MLH1 and its dimer PMS2) and the detection of microsatellite instability (MSI) using PCR.

Keywords: Lynch syndrome (LS); pancreatic neuroendocrine tumors (P-NETs); immunohistochemistry (IHC); microsatellite instability (MSI); polymerase chain reaction (PCR)

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Introduction

Lynch syndrome (LS), previously known as hereditary nonpolyposis colorectal cancer, is an autosomal dominant condition with incomplete penetrance. It is caused by a mutation in one of the deoxyribonucleic acid (DNA) mismatch repair (MMR) genes which leads to its loss of expression and function. Mutations occur most frequently in the MLH1 and MSH2 genes, followed by MSH6 and then finally PMS2 (1-3).

Colorectal carcinoma (CRC) is the neoplasm most commonly associated with LS. It is often diagnosed in young people under the age of 50 years, and accounts for 3–5% of all CRC (1,2). Endometrial carcinoma (EC) is the second most frequent malignancy in women. There are other less commonly associated neoplasms, like ductal adenocarcinomas of pancreas.

Pancreatic neuroendocrine tumors (P-NETs) are the second most common group of pancreatic tumors in the general population (4). Although most P-NETs are sporadic, some of them appear in the context of hereditary syndromes. The most commonly associated syndromes are multiple endocrine neoplastic disease type 1 (MEN1), Von Hippel-Lindau syndrome (VHL), neurofibromatosis type 1 (NF1) and tuberous sclerosis (TS).

Non-functioning pancreatic neuroendocrine tumors (NFP-NET) are the most frequent P-NETs. The coexistence of LS and P-NET has only rarely been described but the association between them has not been confirmed (3).
We report a patient with confirmed LS who developed many of the neoplasms included in the spectrum of this syndrome. We pay special attention to the presence of NFP-NET, in which we demonstrate the relationship between LS and P-NET by immunohistochemistry (IHC) (loss of expression of proteins MLH1 and its dimer PMS2) and the detection of microsatellite instability (MSI) using polymerase chain reaction (PCR).

**Case presentation**

Our patient is a 65-year-old female from a family affected by LS. The index case was a paternal cousin of the patient. Family history is available in Figure 1.

At the age of 45, a regular screening colonoscopy for CRC (due to father’s death related to CRC) revealed left CRC. A sigmoidectomy was performed. The pathology study showed a pT1N0M0 adenocarcinoma. At age 57, she was diagnosed of a right CRC, and a right hemicolectomy was performed. Pathology study confirmed a pT1N0M0.

At age 58, she underwent total abdominal hysterectomy with bilateral salpingo-oophorectomy for uterine cancer. Pathology study showed a pT1aN0M0 low grade endometrioid adenocarcinoma. Given the known family history with a cousin and a brother with LS, MMR protein IHC was performed, which revealed loss of expression of MLH1 and PMS2 proteins in tumor cells of endometrioid carcinoma and both CRC.

As the patient met the Amsterdam II and Bethesda criteria (Table 1), a genetic study was carried out which detected a MLH1 MMR gene mutation, c.731G > A (p.Gly244Asp), in accordance with the mutation found in the affected relatives. The patient’s sons also underwent genetic study but were not carriers of the mutation.

At age 63, a tumor in the right breast was detected on screening mammography and was treated with a lumpectomy. Histology study revealed an invasive low grade ductal carcinoma.

At age 65, a screening gastroscopy (due to the death of a brother with gastric cancer) revealed a duodenal adenocarcinoma. A pancreaticoduodenectomy with pylorus preservation operation was performed. The histology study reported three neoplasms: a pT1N0 duodenal low grade adenocarcinoma, and two incidental NFP-NET in the pancreatic head, one of 11 mm (G2, Ki67: 3%) and another of 7 mm (G1, Ki67: <1%).

A few months later, excision of a skin lesion was diagnosed as sebomatiscoma.

IHC study was carried out on all the tumors presented, which confirmed the lack of nuclear expression of MLH1 and PMS2 proteins and the persistence of MSH2 and MSH6 expression in all but the G1 NFP-NET, in which the four proteins remained expressed (Figures 2, 3).

MSI was also studied. MSI is generated at DNA level due to incorrect functioning of the DNA MMR system. We analyzed five microsatellites (repetitive DNA composed of...
Table 1 Amsterdam II criteria (5) and Bethesda criteria (6)

<table>
<thead>
<tr>
<th>Amsterdam II criteria (all criteria must be met)</th>
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<tr>
<td>Three or more relatives with histologically confirmed colorectal cancer or cancer of the endometrium, small bowel, ureter, or renal pelvis, one affected relative being first-degree relative of the other two; FAP should be excluded</td>
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<tr>
<td>Two or more successive generations are affected</td>
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<td>At least one relative was diagnosed before the age of 50 years</td>
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<th>Revised Bethesda guidelines (one or more of the following criteria must be met)</th>
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<tr>
<td>Colorectal cancer before the age of 50 years</td>
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<tr>
<td>Synchronous or metachronous colorectal cancer or other HNPCC-related tumors, regardless of age</td>
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<td>Colorectal cancer with MSI-H morphology before the age of 60 years</td>
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<tr>
<td>Colorectal cancer (regardless of age) and a first-degree relative with colorectal cancer or an HNPCC-related tumor before the age of 50 years</td>
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<tr>
<td>Colorectal cancer (regardless of age) and two or more first- or second-degree relatives diagnosed with colorectal cancer or an HNPCC-related tumor (regardless of age)</td>
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FAP, familial adenomatous polyposis; MSI, microsatellite instability.

Figure 2 IHC studies (10x) of duodenal adenocarcinoma. (A) Nuclear loss of expression for MLH1 protein; (B) nuclear loss of expression for PMS2 protein; (C) MSH2 protein nuclear expression; (D) MSH6 protein nuclear expression. Neoplastic gland structures can be identified infiltrating the duodenal submucosal layer. Gland nuclei show loss of MLH1 and PMS2 expression (A,B) and maintain MSH2 and MSH6 expression (C,D). IHC, immunohistochemistry.
sequences ranging in length from one to five DNA base-pairs) which are monomorphic and highly stable in the normal population. If the result shows two or more of them with extra DNA sequences, it is considered unstable (MSI-H) and is attributed to a dysfunction in DNA repair. If additional sequences are not found in any microsatellite, it is considered stable (MSI-S). Occasional cases show additional sequences in only one microsatellite, and it is considered as low unstable (MSI-L).

The study was conducted by PCR on patient samples, which showed MSI-H in colonic, duodenal and skin neoplasms, but were stable in the other samples (Figure 4).

Follow-up of patient showed no recurrences three years after the last surgery.

**Discussion**

This is the first case described in the literature that associates LS and NFP-NET. LS is caused by inactivating mutations of DNA MMR genes that impair DNA MMR mechanisms (1-3).

The diagnosis of LS is based on clinical suspicion (Table 1) and confirmed by germline genetic study of mutations in MMR genes (in order of frequency: MLH1, MSH2, MSH6 and PMS2) (1,2,5,6). LS is associated with CRC at an early age, and EC in female. Ovarian, gastric, urothelial, intestinal, pancreatic, hepatobiliary and breast carcinomas are also associated with LS, as well as skin and central nervous system tumors (1,2,6). Nowadays, families comprise fewer members than some decades ago, so there is less chance to diagnose LS based on clinical criteria.

IHC study of tumor specimens allows detection of the loss of expression of proteins encoded by MMR genes (1,3). In the case of an MLH1 gene mutation, loss of expression of the encoded protein prevents protein dimer formation with PMS2, which is recruited after MLH1. In our case, loss of expression of PMS2 is conditioned by the mutation detected in MLH1. This feature was observed by IHC in all the tumors analyzed with the exception of NFP-NET G1 (Figures 2,3).

MSI study of tumor samples was performed too (3). MSI-H was identified in colon, duodenum and skin neoplasms, but not in the others (Figure 4). This may have been because the microsatellites studied to detect MSI-H by PCR in our laboratory were more specific for CCR, so the absence of MSI-S detection in other malignancies does not rule out the diagnosis of LS. Thus, loss of protein expression, even in the case of a negative PCR result, is

![Figure 3 IHC studies (10x) of G2 NFP-NET. (A) Nuclear loss of expression for MLH1 protein; (B) nuclear loss of expression for PMS2 protein; (C) MSH2 protein nuclear expression; (D) MSH6 protein nuclear expression. Neoplastic cells display a trabecular pattern. Tumor cell nuclei show loss of MLH1 and PMS2 expression (A,B) and maintain MSH2 and MSH6 expression (C,D). IHC, immunohistochemistry.](image-url)
Some data have associated LS and pancreatic tumors. There is a cumulative risk of pancreatic adenocarcinoma in LS patients of 1.31% (95% CI: 0.31–2.32%) at 50 years and 3.68% (95% CI: 4.7–15.7%) at 70 years compared to the general population (8). This association is included in the latest review of the Bethesda Guidelines (6).

Most pancreatic adenocarcinomas described as associated with LS reveal mutations in the MMR repair gene MSH2 (7,9,10), as often happens in extra-colonic tumors. Kastrinos et al. report that of the 47 cases of pancreatic adenocarcinomas, 31 had a mutation in the MSH2 gene, 13 cases in the MLH1 gene and three in the MSH6 gene (7). The most common pancreatic tumors associated with LS are ductal adenocarcinoma (7,10), although some cases of intraductal papillary mucinous neoplasm (IPMN) have been suggested (9).

P-NET is the second most frequent pancreatic tumors. The non-functioning type is the most prevalent (4). Clinical manifestations are usually silent or poor, often diagnosed incidentally or because of a mass effect. Although most P-NET is sporadic, some appear in the context of a hereditary syndrome, the most common are: MEN1 disease, VHL disease, NF1 and TS.

Karamurzin et al. described a case of NFP-NET in a patient with LS, but IHC examination of tumor did not show loss of expression of any MMR protein and there were no MSI-H (3). Therefore, the association between LS and P-NET has been suggested but has not been confirmed to date (3).

In our patient, a germline mutation of MLH1 was demonstrated in the biggest NFP-NET. But it was not demonstrated in the little one. Hypothesis for the retained protein expression in the small one is that complete loss of DNA, deoxyribonucleic acid.
MMR function requires multiple cell cycle alterations and could be a late event in the neoplastic transformation (9).

Small intestine tumors account for fewer than 2% of gastrointestinal tract tumors in the general population (11). There are several inherited syndromes that increase the likelihood of developing this condition, such as familial adenomatous polyposis (FAP) disease, Peutz-Jeghers disease and LS (11,12). In LS, the probability of small intestine carcinoma ranges from 1–4%, around 100 times higher than in the general population (11,12). Adenocarcinomas are the most common type (as in our patient), with one reported case of carcinoid-type neuroendocrine tumor. Involvement of duodenum and jejunum is higher than terminal ileum. Attempts to determine whether the risk of small intestine carcinoma differs according to the MMR protein affected have shown it to be higher in cases of MLH1 and MSH2 mutations, with no difference between them (11). Incidence of small intestine carcinoma is so low that screening tests such as capsule endoscopy or gastroscopy are not recommended (13). Nevertheless, an increased detection of duodenal tumors has been observed with gastroscopy explorations performed as screening for the detection of gastric carcinoma (14). We consider that gastroduodenoscopy should be indicated in patients with LS as a screening test for gastroenteric neoplasms.

Conclusions

Our report supports the inclusion of P-NET in the list of extracolonic LS-associated tumors. P-NET must be included in the screening list as well as duodenal adenocarcinomas. We consider that in LS patients, a gastroduodenoscopy with endoscopic ultrasonography should be considered and an enhanced CT scan with an arterial and portal vein phase could help to discover P-NET tumors.

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Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

Informed Consent: Written informed consent was obtained from the patient for publication of this manuscript and any accompanying images.

References


