INTRODUCTION

Neuroendocrine neoplasms account for 1.2-1.5% of all gastrointestinal neoplasms, with an annual incidence of 1.6-2.0 new cases for every 100000 persons. These neoplasms may involve many different organs and sites, with a majority of the tumors occurring in small bowel, appendix, rectum, and the stomach (1). Biliary neuroendocrine neoplasms are extremely rare, and represents only 1% of all gastrointestinal neuroendocrine neoplasms. This is probably due to a very low numbers of neuroendocrine (Kulchitsky) cells in the biliary epithelium (2, 3). Because of its extremely rare incidence and presenting symptoms like painless jaundice, most extrahepatic biliary neuroendocrine neoplasms are misdiagnosed as adenocarcinoma prior to surgery (4). Most studies on these tumors consist of case reports with literature review and focus on histologic features or clinical outcomes.

This case is unique in that the role of imaging study in the diagnosis of neuroendocrine neoplasm of biliary duct has not been reported until now.

CASE REPORT

A 64-year-old man with a known diagnosis of pneumoconiosis was transferred to our hospital due to indigestion and jaundice. The initial abdominal CT and MRI revealed a 2.0 cm enhancing mass in the proximal common bile duct (CBD) with several enlarged lymph nodes. The mass was presumed to be a cholangiocarcinoma, and a CBD segmental resection and choledochojjunostomy was performed. However, the final diagnosis was that of a mixed endocrine-exocrine carcinoma, a high-grade neuroendocrine neoplasm. Seven months after the operation, a follow-up abdominal CT study revealed multiple small arterial enhancing nodules in both hepatic lobes. A sono-guided liver biopsy confirmed these as metastatic mixed endocrine-exocrine carcinoma. This case is unique in that the imaging study regarding the neuroendocrine neoplasm of biliary duct has not been previously reported.

Index terms
Neuroendocrine Neoplasm
Mixed Adenoneuroendocrine Carcinoma
Biliary Duct
World Health Organization Classification
Imaging Findings of Neuroendocrine Neoplasm in Biliary Duct with Liver Metastasis

A well-enhancing 2.0 cm polypoid mass in the proximal common bile duct (CBD), which resulted in extrahepatic biliary obstruction. The mass demonstrated about 100 Hounsfield unit (HU) net enhancement (17 HU on pre-enhancement image, 113 HU on post-enhancement image) during the late arterial phase. Several enlarged lymph nodes were seen around the common bile duct, porta hepatitis, and paraaortic space. Seven days after the CT scan, the patient underwent percutaneous transhepatic biliary drainage (PTBD) procedure. The percutaneous tubography revealed a small expansile intraductal polypoid mass in the proximal CBD with moderate dilatation of the intrahepatic ducts (Fig. 1C). Positron emission tomography (PET) CT showed a faint fluorodeoxyglucose (FDG) uptake [standardized uptake value (SUV) max: 2.4] in this mass.

Seven days subsequent to PTBD, CBD segmental resection and choledochojunostomy was performed under a presumed diagnosis of cholangiocarcinoma. Gross examination revealed a 2.0 x 2.0 cm polypoid mass. Microscopically, the tumor had invaded beyond the wall of CBD to the surrounding adipose tissue with one regional lymph node metastasis. Histological examination (Fig. 1D, E) revealed the tumor to have both adenocarcinomatous component and neuroendocrine tumor. Immunohistochemistry was positive for COX-2, suggestive of adenomatous component, and also revealed positive expression of CD56, chromogranin, neuron-specific enolase, suggesting neuroendocrine component. The Ki67 labeling index, a proliferative marker, was 95%. Thus, the final diagnosis was confirmed as mixed endocrine-exocrine carcinoma.

The postoperative course was uneventful, until a follow-up abdominal CT (Fig. 1F, G) at 7 months revealed multiple nodules in both hepatic lobes. These lesions were small, arterial enhancing, and demonstrated subtle delayed washout. This imaging study was suggestive of hepatocellular carcinomas. On ultrasonographic exam, these masses were homogenous hypere-
choic nodules in the liver parenchyma. Sono-guided liver biopsy was performed for several lesions. The pathology was confirmed to be metastatic mixed endocrine-exocrine carcinoma. The patient is currently alive undergoing chemotherapy treatment.

DISCUSSION

On one hand, biliary neuroendocrine neoplasms are extremely rare and represent only 1% of all gastrointestinal neuroendocrine neoplasms. On the other hand, adenocarcinoma is the most common type of all biliary duct tumors, accounting for approximately 80% of occurrence (2, 5). The most common presenting symptom of biliary neuroendocrine neoplasms is jaundice, and the most common sites are the common bile duct (58%), perihilar region (28%), cystic duct (11%), and the common hepatic duct (3%) (3). Most biliary neuroendocrine neoplasms are misdiagnosed before as adenocarcinoma preoperatively because of its extremely rare incidence and its similarity to the presenting symptoms and nonspecific radiologic findings of adenocarcinoma (4). The presenting symptoms involve jaundice, biliary colic pain, abdominal discomfort, and weight loss. Unlike those found in other locations, neuroendocrine neoplasms of the extrahepatic biliary ducts tend to have an indolent biological behavior, even when metastatic. This makes it more difficult for a correct diagnosis. The variety of nomenclature used for neuroendocrine neoplasm of the gastrointestinal tract add to the confusion among clinicians and radiologists.

Recently, a new classification of neuroendocrine neoplasm has been proposed and adopted by the World Health Organization (WHO). The WHO 2010 classification identified following histopathologic subtypes: 1) neuroendocrine tumor (NET) G1 and G2, which is defined as a well differentiated neuroendocrine neoplasm; 2) neuroendocrine carcinoma (NEC), which is a poorly differentiated, high grade malignant neoplasm; and, 3) mixed adenoneuroendocrine (exocrine endocrine) carcinoma, which has a morphologic phenotype of both gland forming epithelial and neuroendocrine tumor cells and is defined as a carcinoma because both components are malignant at least 30% of all cases.

Currently, many diagnostic imaging tools are available for biliary tumors, including abdominal ultrasonography, CT, and MRI. Because of the nonspecific nature of radiologic findings, the final diagnosis of neuroendocrine neoplasms is usually made postoperatively with histological and immunohistochemical examination of the surgical specimen (6). No reports in the radiology literature described the appearance of biliary neuroendocrine neoplasms. As a result, the assessment of imaging features of those tumors had been limited. CT findings of non-biliary neuroendocrine neoplasms vary with size and location. However, biliary neuroendocrine neoplasms are generally more prominent in the arterial phase and less prominent in the portal venous and equilibrium phases, as are most hypervascular tumors. In some case reports, most (6/7; 85.7%) biliary neuroendocrine neoplasms had a polyloid appearance. The mean diameter of these tumors was 2.94 cm (2-4.5 cm) (3, 5-8). Generally, neuroendocrine tumor demonstrate lower 18F-FDG uptake on a PET CT. However, this finding is dependent on the neoplasm grade, as high grade neuroendocrine neoplasms have significantly higher uptake of 18F-FDG, when compared with that of low grade neuroendocrine neoplasms (median SUV 11.7 vs. 2.9). Nevertheless, 18F-FDG uptake was not seen in few cases of high grade neuroendocrine neoplasms (9).

In this study, contrast enhanced CT images revealed a 2 cm polypoid mass in the proximal CBD with regional lymph node (LN) involvement. This proximal CBD mass had a relatively marked enhancement during the late arterial phase, which was in contrast to the usual findings for cholangiocarcinomas. The PET CT demonstrated faint 18F-FDG uptake (SUV max: 2.4) in the mass. This finding may be because the tumor was too small for significant 18F-FDG uptake or because of the wide spectrum of neuroendocrine neoplasms displaying variable amount of 18F-FDG uptake.

Biliary neuroendocrine neoplasms are metastasize more frequently to the liver, pancreas, gallbladder, and in the regional lymph nodes (6, 10). According to the study by Kim et al. (10), biliary neuroendocrine neoplasms metastasized to hepatic parenchyma in 2/7 cases of pure neuroendocrine carcinoma and in 2/7 cases of mixed endocrine-exocrine carcinoma. In contrast, hepatic metastasis was not reported for NET G1 and G2 (0/6) tumors. LN metastasis of mixed endocrine-exocrine carcinoma (6/7; 85.7%) was distinguished from others (5/13; 38.5%). The results of that study corresponds with our case. According to the histopathologic subtypes in the WHO 2010 classification, biliary neuroendocrine neoplasms had a wide spectrum of clini-
cal outcomes, including differences in recurrence rates, disease free survival, and overall survival (10).

In conclusion, the characteristic imaging findings of biliary neuroendocrine neoplasms are relatively marked enhancement during the arterial phase, polyloid shape, small tumor size (2-4.5 cm), and generally lower uptake of $^{18}$F-FDG depending on tumor grade. The existence of hepatic or LN metastasis in neuroendocrine neoplasms raises a possibility of high grade neuroendocrine neoplasms, such as NEC or mixed endocrine-exocrine carcinoma, as was our case. The imaging study results for extrahepatic cholangiocarcinomas may differ from those of biliary neuroendocrine neoplasms. Infiltrating extrahepatic cholangiocarcinoma is the most common type, and it manifests as a high-attenuation mass encircling the lumen or thickened wall at the site of biliary obstruction. Polypoid extrahepatic cholangiocarcinoma is the second most common type of extrahepatic cholangiocarcinomas. It manifests as a low-attenuation mass within the dilated bile duct and frequently demonstrates extensive superficial spreading, resulting in diffuse involvement (11).

Our report is valuable in that it is the first case report with a focus on imaging study findings of neuroendocrine neoplasms of biliary duct with liver metastasis, the correlations between histopathologic and these radiographic findings in the context of the newer WHO classification.

REFERENCES

간전이를 동반한 담관내 신경내분비종양의 영상소견: 중례 보고

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64세 남자 환자가 소화불량과 황달을 주소로 우리병원에 전원되었다. 내원시 전산화단층촬영과 자기공명영상 검사에서 근위부 총담관에 약 2 cm 크기의 조영증강되는 종괴가 발견되었고, 여러 개의 거친 부위 림프절을 동반하였다. 종괴는 담관암으로 생각되었으며 따라서 총담관의 구역절제술과 총담관공장연결술을 시행하였다. 그러나 종괴는 신경내분비 종양의 일종인 혼합형 내분비-외분비 암종으로 진단되었다. 7개월 후에, 추적검사로 시행한 복부 전산화단층촬영에서 양측 간엽에 여러 개의 작은 결절들이 보였고 이들은 동맥기에 조영증강을 보였다. 초음파 유도하 간생검에서 이는 혼합형 내분비-외분비 암종으로 진단되었다. 이것은 지금까지 보고된 적이 없었던 담관내 발생한 신경내분비종양의 영상소견에 대한 증례이다.

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