Well-Differentiated Mucinous Cystadenocarcinoma of the Renal Pelvis

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Abstract

Adenocarcinoma arising from renal pelvis is uncommon. A mucinous cystadenoma of the renal pelvis with malignant transformation is a rare entity. We report a case of well-differentiated mucinous cystadenocarcinoma of the renal pelvis with malignant transformation of cystadenoma with mucus-filled kidney (muconephrosis) due to mucin produced by the tumor.

Key words: Mucinous adenocarcinoma, malignant transformation, cystadenoma, renal pelvis

Case Report

Introduction

A cystadenoma or cystadenocarcinoma of renal pelvis is a rare neoplasm. A mucinous cystadenoma of the renal pelvis with malignant transformation is a relatively rare entity. Adenocarcinoma accounts for less than 1% of malignancies arising from renal pelvic epithelium. Spires et al. subclassified these tumors into three histologic types: tubulovillous, mucinous and papillary nonintestinal. Most reported cases are malignant (1-5). We report a case of well-differentiated mucinous cystadenocarcinoma of the renal pelvis with malignant transformation of cystadenoma. Mucus-filled kidney (muconephrosis) due to mucin produced by the tumor was detected.

Case

A 67-year-old man presented with intermittent left flank pain during ten months and had swollen abdomen. There had been no other diagnosed malignancy especially about gastrointestinal system in his history before. The clinical diagnosis was hydronephrosis due to a cystic lesion as hydatid cyst. Left simple nephrectomy was performed with maintaining the integrity of cyst. The gross examination of the specimen revealed that this mucoid fluid was connected with cyst wall. There was only a huge unilocular cystic lesion measured 20x17.5x8 cm. Renal medulla, cortex and pelvicaliceal system were not obviously observed (Figure 1). The thickness of cyst wall was 0.1-1.5 cm. 2.7 cm diameter mucinous mass with villous structures was detected in an area of cyst lumina. No calculi were found. On histology, the cyst wall showed single or generally pseudostratified-layered, tall columnar mucinous epithelium with some goblet cells as intestinal epithelium. In the fibrocollagenous stroma, there were hemorrhage, necrosis, inflammatory infiltrate and acellular mucin.
Mucinous cystadenocarcinoma of renal pelvis

Figure 1. The huge, unilocular, mucinous cyst. Renal medulla, cortex and pelvicaliceal system are not obviously observed.

Figure 2. H&E X400-The cyst wall has pseudostratified-layers, tall columnar mucinous epithelium with nuclear atypia which formed villous structures.

Figure 3. The epithelium shows immunohistochemical Cytokeratin 7 expression.

Figure 4. Immunohistochemical staining shows a positivity for Cytokeratin 20.

DISCUSSION

Adenocarcinoma arising from renal pelvis was first reported by Grohe in 1901 (6) and in 1973 Aufderheide and Streitz reported the first major review (2). Spires et al. reviewed 59 cases in 1993 which is the most recent review (4). Chronic infection, renal calculi and hydronephrosis were involved as possible aetiological factors in mucinous adenocarcinoma of the renal pelvis (4,7-8). The normal transitional epithelium can transform squamous, columnar, or cuboidal form with metaplasia (9). Conversion of transitional epithelium to colonic type epithelium is relatively rare. Gangane et al. be-
lieve that mucinous cystadenoma, like its malignant counterpart, arises from foci of intestinal metaplasia, and say that they were able to demonstrate histological transition from transitional epithelium to metaplastic mucinous areas in the same lesion (10). However, Park et al. reported a case of renal pelvic villous adenoma and diffuse intestinal metaplasia of the urothelium without gastrointestinal pathological findings (11). Kobayashi et al. have related the histogenesis adenocarcinoma to metaplasia of transitional cell carcinoma (7). There have been cases as mucinous cystadenoma of the renal pelvis with malignant transformation (2) as in our case, a mucin-producing cystadenoma, borderline malignancy of the renal pelvis and ureter (1), mucinous cystadenoma in a horseshoe kidney (12,13), intestinal type of villous adenoma (14) and mucinous adenocarcinoma of renal pelvis and villous adenoma of bladder after caecal augmentation of bladder (15).

Our case is a mucin-producing cystadenoma with malignant transformation without a history of long-standing chronic infection and renal stone, also there was no enteral communication and gastrointestinal pathology. There was hydronephrosis due to mucin secretion of tumor (muconephrosis). We have used the same criteria used for mucinous ovarian tumors for the diagnosis of malignancy. Ross and D’Amato (13) suggested that it might also be appropriate for mucinous tumors of the kidney.

REFERENCES