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A CASE LARGE PROTRUDING CYSTITIS GLANDULARIS: CLINICAL, HISTOLOGICAL AND MUCIN-HISTOCHEMICAL STUDY

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A case of large protruding cystitis glandularis is reported. A 36-year-old man was admitted to our hospital due to acute cholecystitis, and large protruding masses were incidentally found in the urinary bladder by abdominal ultrasonography. The histological study revealed that they consisted of a large number of Brunn's nests with or without cysts which were often accompanied with columnar epithelial metaplasia, and of glandular structures closely resembling the colonic crypts. The mucin-histochemical study demonstrated glandular lesions in the bladder secreted colonic type mucin, and endocrine cells positive with Grimelius' staining. A review of literature disclosed 19 clinical cases of cystitis glandularis, since 1970, in Japan, but such a large protruding lesion as this case is rare. We first performed detailed histological and mucin-histochemical studies for this clinical case.

Key words: Cystitis glandularis, Protruding lesion, Mucin-histochemistry

INTRODUCTION

Cystitis glandularis is one of the proliferative lesions which are commonly found in the urinary bladder, but it does not usually show a large protruding lesion in the urinary bladder. Presence of mucin-secreting glandular structures, closely resembling colonic crypts, is well recognized as a metaplastic change.

We experienced a case of large protruding cystitis glandularis, and performed detailed histological and mucin-histochemical studies.

CASE REPORT

A 36-year-old man was admitted to the Department of Internal Medicine of our hospital in October 1988 due to acute cholecystitis. By abdominal ultrasonography, large protruding masses were incidentally found in the urinary bladder. After recovery from acute cholecystitis, he was transferred to the Department of Urology for the purpose of detailed examinations and treatment for the vesical lesion on November 4, 1988. He had no urinary symptoms such as burning and difficulty on urination or hematuria.

Physical, hematological, and blood chemical examinations were within the normal limit. Urinalysis was normal and urinary cytology was negative. He had no difficulty in urination and no vesicoureteral reflux (VUR).

Excretory urography showed no hydro-nephrosis, but slight dilatation was seen in the lower portion of both ureters. On the urethrocystogram several filling defects of the urinary bladder were noted. CT scan demonstrated sessile protruding masses from the base of the bladder, but did not show any finding suggesting extravesical invasion. Enhanced CT scan in the prone position revealed the tip of the protruded mass enhanced strongly (Fig. 1).

Cystoscopy revealed several finger-sized nodular and sessile masses at the bladder neck and trigone, part of which showed a tower or a beak-like appearance (Fig. 2).
Enhanced CT scan demonstrates a sessile protruding mass from the base of the urinary bladder and its tip is enhanced.

Cystoscopy reveals some finger-sized sessile masses with a tower or beak-like appearance at the neck and trigone of urinary bladder.

The left ureteral orifice was not recognized due to the tumorous lesion. Biopsy of this lesion disclosed a cystitis glandularis with colonic-type gland formations. Transurethral resection of this benign lesion was performed on November 16, 1988. The border of the lesion was not so clearly identified.

Eight months after the first operation cystoscopy disclosed recurrence of similar masses at the trigone, and transurethral resection was performed again. Histopathological diagnosis was the same as the previous one without any malignant change. Six months after the second operation, cystoscopy disclosed recurrence at the trigone. Transurethral resection was performed for the third time.

**PATHOLOGY**

The tissue specimens were fixed in 10% formalin for 24 hours and embedded in paraffin. In addition to a routine hematoxylin and eosin staining, to clarify the exact character of the mucin secreted, we applied several specific mucin stainings as follows: (1) periodic acid-Schiff staining (PAS), (2) high iron diamine and alcian blue (pH 2.5) stainings (HID-AB), (3) periodic acid-borohydride, potassium hydroxide and PAS stainings (PB-KOH-PAS), (4) paradoxical concanavalin A for stable class III stainings (con A (III)). PAS stains non-specifically various kinds of mucin. By HID-AB, the mucin blue in color is sialomucin, and that black in color is sulphomucin. The mucin of the small intestine and colon is positive with HID-AB. PB-KOH-PAS indicated the presence of O-acetylated sialomucin, specific to the goblet cells of the colon. Con A (III) is specific to the gastric type mucin. We also applied Grimelius’ staining to find endocrine cells.

Histologically, in the lamina propria of the urinary bladder, there were many Brunn’s nests (Fig. 3a) with or without cysts, which were often accompanied with columnar epithelial metaplasia (Fig. 3b). Glandular structures, composed of goblet cells, were also seen to be scattered. They closely resembled colonic crypts (Fig. 3c). Neither adenomatous nor carcinomatous

**Fig. 3.** a. Brunn’s nests, b. Brunn’s nests with cyst lined by columnar epithelial cells, c. Glands resembling colonic glands (H.E. stain)
findings were seen in these cells. These goblet cells were positive with PAS, HID-AB, and PB-KOH-PAS (Fig. 4a). In HID-AB cells with sialomucin were nearly equal to those with sulphomucin. However, they were negative with con A (III). Brunn’s nests were negative with PAS, HID-AB, PB-KOH-PAS and con A (III). Columnar epithelial metaplasia in Brunn’s nests was slightly positive with PAS and HID-AB, but it was negative with PB-KOH-PAS and con A (III). Cells positive with Grimelius’ staining were seen to be scattered in the glands (Fig. 4b) and in the Brunn’s nests.

**DISCUSSION**

Cystitis glandularis is one of the proliferative lesions commonly found in the urinary bladder. Ito et al. described that cystitis glandularis was found in 71% of the 125 autopsy cases of macroscopically normal-appearing urinary bladders of both sexes and all ages. On the other hand, including the present case, only 20 cases of cystitis glandularis with clinical symptoms or findings have been reported since 1970 in Japan. Age distribution is from 6 to 80 years old with the average of 46 years, and it is more frequently seen in males than in females. Irritative symptoms of the bladder are most frequent and some have gross hematuria or difficulty in urination. The incidence of these symptoms is high in the bladder neck and trigone.

In Japan, detailed histological studies have not been reported for the clinical cases. Koss described that cystitis glandularis cannot be clearly differentiated from cystitis cystica. However, in cystitis glandularis, cells lining the glands may be cuboidal and resemble colonic epithelium with goblet cells. According to WHO, this lesion is classified as glandular metaplasia (glandular “cystitis”), which is characterized by mucus containing columnar epithelial cells either on the surface or forming glands in the lamina propria.

Many authors have reported that these mucus containing cells are positive with PAS. However, PAS non-specifically stains various kinds of mucin. Therefore, we applied detailed mucin-staining as described above, to clarify the property of the mucin in the present case.

In this study, O-acetylated sialomucin was demonstrated in the goblet cells. Therefore, mucin of the goblet cells in cystitis glandularis was the same as that of the colon. Recently, Hamid et al. also demonstrated endocrine differentiation in inflamed urinary bladder epithelium with a metaplastic change. By Grimelius’ staining, we also found endocrine cells both in the colonic-type glands and Brunn’s nests. Therefore, this case also demonstrated that the bladder mucosa has a potential to change toward a more differentiated glandular structure of colonic type, via Brunn’s nests, as a metaplastic change.

Hasegawa et al. reported using the same method as ours that the colonic type mucin was demonstrated not only in cystitis glandularis in autopsy cases but also in half of the cases of mucin-producing adenocarcinomas of the bladder. Wells et al. reported that adenocarcinoma of the
bladder associated with cystitis glandularis produced O-acetylated sialomucin, whereas a primary adenocarcinoma of urachal origin did not. Thus, they showed that PB-KOH-PAS may be used to differentiate primary adenocarcinomas from primary urachal carcinomas of the urinary bladder.

These findings are not only very interesting to explain the histogenesis of the urinary bladder, but also suggest that cystitis glandularis is related to the development of adenocarcinoma in the urinary bladder. There are several reports on cystitis glandularis which underwent a malignant change during follow-up period of 5~15 years. On the other hand, it is evident, as systematically demonstrated by Ito et al., that cystitis glandularis is a very common autopsy finding. Clinically, Ohigashi et al. reported a case of cystitis glandularis followed for over 10 years without malignant changes.

As a clinical practice, the patient with cystitis glandularis must be placed on the risk chart, and prolonged close follow up is obligatory.

REFERENCES


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大きな腫瘍を形成した腺性膀胱炎の１例：臨床的・組織学的・細胞組織化学的検討

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大きな腫瘍を形成した腺性膀胱炎の１例を報告する。症例は36歳男性で急性膀胱炎にて当院入院中、腹部超音波検査にて偶然に膀胱内に腫瘍が発見された。組織学検査の結果、部分的に円柱上皮化生を伴った囊胞をもつ、多くのブラウン炎症、結腸上皮にきわめて類似した多くの腺構造が認められた。粘液組織化学染色の結果、これらの腺は結腸型の粘液を分泌しており、またグリセリウス染色陽性の分泌細胞も認められた。本邦では1970年より今日までに19例の腺性膀胱炎の臨床例が報告されているが、本例のように大きな腫瘍を形成する例は少ない。また、臨床例に対して組織学的・細胞組織化学的に詳しい検討を加えたのは本例が初めてである。

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