INTRODUCTION

Desmoid tumors are rare benign tumors with aggressive fibroblastic proliferation. Although desmoid tumors do not metastasize, they have locally aggressive features and can cause a urinary fistula. Here, we report a case of a 35-year-old woman with Gardner syndrome who was diagnosed with an intra-abdominal desmoid tumor 1 year previously and who presented with a newly developed cystic mass lesion on a computed tomography scan. The cystic mass lesion was clinically diagnosed as an urinoma from the right ureterotumoral fistula; thus, surgical resection of the mass lesion was planned. However, Tc-99m diethylenetriamine pentaacetic acid renal scintigraphy revealed bilateral ureterotumoral fistulas; hence, the treatment plan was changed to conservative management.

Index terms
Desmoid Tumor
Radionuclide Imaging
Urinary Fistula
Diethylenetriamine Pentaacetic Acid

CASE REPORT

A 35-year-old woman with Gardner syndrome presented to our medical center with abrupt onset abdominal pain. She had undergone total colectomy at the age of 29 years, and a mass lesion in the small bowel mesentery detected 1 year previously was histologically confirmed to be a desmoid tumor (Fig. 1). She was subsequently treated with several chemotherapy cycles for the desmoid tumor, but the tumor did not respond. She had undergone a right percutaneous nephrostomy (PCN) during chemotherapy due to hydronephrosis caused by the desmoid tumor.
The patient's vital signs were stable on admission. Laboratory results revealed the upper normal limit of the white blood cell (WBC) count (10.2 × 10^3/μL; normal range, 4.0–10.8 × 10^3/μL), a low hemoglobin level (9.7 g/dL; normal range, 12.0–16.0 g/dL), a low platelet count (100.0 × 10^3/μL; normal range, 130–400 × 10^3/μL), a normal creatinine level (1.2 mg/dL; normal range, 0.5–1.2 mg/dL), and a high C-reactive protein level (69.8 mg/dL). The results of a WBC differential test revealed a high differential neutrophil count of 92.2%.

Contrast-enhanced abdominal computed tomography (CT) was performed to evaluate the abdominal pain, which revealed the already identified desmoid tumor in the small bowel mesentery but no significant interval change in its size compared to that on the previous CT scan. However, a new 5 cm in size poorly-enhanced infiltrative mass lesion containing fluid and a small air density was detected at the mesenteric root (Fig. 1). An abscess combined with the desmoid tumor was suspected. Percutaneous drainage (PCD) of the new cystic mass lesion was performed. After PCD and antibiotic treatment, the cystic mass lesion shrank, but it continued to drain. The color of the leaking fluid was clear and yellowish, suggesting urine. No definite connection was observed between the ureter and the new mass lesion on CT, but a right ureterotumoral fistula was suspected based on the location of the mass. We suspected a right ureterotumoral fistula and an urinoma from urine leakage based on the CT findings and planned to surgically resect the mass and urinoma and repair the right ureter.

Tc-99m DTPA renal scintigraphy was performed preoperatively to assess the ureterotumoral fistula. Cut-off of the radioactivity was observed at bilateral mid-ureters during the excretion phase on dynamic renal scintigraphy images, suggesting bilateral ureterotumoral fistulas (Fig. 2). No radioactivity was observed in the cystic mass lesion because of PCD. On the other hand, radioactivity was found in the urine contained in the PCD tubes and the PCN tubes and urinary bladder, confirming the presence of an ureterotumoral fistula. F-18 fluorodeoxyglucose (FDG) positron emission tomography (PET)/CT was also performed to detect a hidden malignancy other than the desmoid tumor (Fig. 3). The PCD tube was locked just before the PET/CT scan and urine activity was seen instantly inside the cystic mass lesion. No abnormal increase in FDG uptake was detected to suggest a hidden malignancy, and the desmoid tumor lesion only showed a mildly increased FDG uptake.

As unexpected bilateral ureterotumoral fistulas were detected on renal scintigraphy, removal of the tumor could result in bilateral ureteral dysfunction. Therefore, the treatment plan was changed from surgical resection to conservative management with medication. The patient has been receiving regular clinical follow-up at our medical center.

**DISCUSSION**

Renal scintigraphy is a very sensitive and non-invasive modality for evaluating renal function and detecting urinary leakage (10). Our patient was clinically suspected to have an ureterotumoral fistula due to the desmoid tumor; however, contrast-en-

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**Fig. 1.** A 35-year-old woman with Gardner syndrome.

A. Transaxial CT images of the patient. A histologically confirmed desmoid tumor is located in the small bowel mesentery on preoperative abdomen CT (arrow).

B. One year later, contrast-enhanced abdominal CT was performed to evaluate the abdominal pain. The already identified desmoid tumor (arrow) is noted in the small bowel mesentery.

C. A new poorly-enhanced cystic mass lesion with an air bubble (arrow) is found in the small bowel mesentery. CT = computed tomography
Ureteral Fistula on Renal Scintigraphy

hanced CT images did not reveal a definite connection between the ureter and the mesenteric mass lesion. As functional nuclear images can detect pathological lesions early before significant anatomical changes, we were able to diagnose the involvement of bilateral ureters by the tumor and change the treatment plan. However, although renal scintigraphy demonstrated bilateral ureterotumoral fistulas, the anatomical location of the fistulas was not clear. Renal scintigraphy using a gamma camera integrated with CT would be more helpful in this case. Hybrid single photon emission computed tomography/CT imaging provides both functional and anatomical information, which demonstrates the superior diagnostic value of functional-anatomical hybrid imaging (5, 6).

The optimal treatment for desmoid tumors is controversial and difficult to determine. Clinical follow-up with close observation is recommended for stable asymptomatic patients with a desmoid tumor (1). However, surgical removal is considered to be the first-line treatment for symptomatic patients, and complete resection of the desmoid tumor with negative microscopic resection margins is the standard goal (1). However, many clinicians are reluctant to perform a surgical resection because of the high recurrence rate. Moreover, the surgical trauma could cause a recurrence by itself (7-9). Complete surgical resection of an in-
tra-abdominal desmoid tumor is extremely difficult and it may be impossible in some cases (1, 7). Because incomplete surgical removal produces a high risk of recurrence and a second operation for treating a recurrent tumor is very difficult to perform due to the intra-abdominal adhesions caused by the previous operation, the surgical approach for an intra-abdominal desmoid tumor should be considered cautiously (2). In addition, patients with familial adenomatous polyposis-associated desmoid tumors have an even higher risk of recurrence (1). Our patient was diagnosed with Gardner syndrome and had had a refractory desmoid tumor with right hydronephrosis. Because of the new symptoms and urine leakage, complete surgical removal of the desmoid tumor and urinoma with repair of the right ureter was preferentially considered, in spite of the deteriorated right ureter function. However, because Tc-99m DTPA renal scintigraphy demonstrated bilateral ureterotumoral fistulas, conservative management was selected to avoid complications that may occur after surgical resection.

We report a case of a patient with an intra-abdominal desmoid tumor who showed unexpected bilateral ureterotumoral fistulas on Tc-99m DTPA renal scintigraphy. Tc-99m DTPA renal scintigraphy was helpful to confirm the diagnosis of an ureterotumoral fistula and to decide the treatment plan.

REFERENCES

신장 신티그래피에서 발견된 요관-종양 샛길을 동반한 
데스모이드 종양 1예

유현우1·이규택2·이상미3*

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데스모이드 종양은 드문 양성 종양으로 침습적인 섬유모세포성 증식을 한다. 데스모이드 종양은 전이되지 않지만, 국소적으로 침윤하는 특성을 보이며 요관-종양 샛길을 형성할 수 있다. 저자들은 1년 전 복부 내 데스모이드 종양으로 진단되었던 가드너증후군의 과거력이 있는 35세 여자 환자에서 컴퓨터단층촬영 후 새롭게 남성 종괴가 발견된 증례를 보고하고자 한다. 남성 종괴는 임상적으로 데스모이드 종양에 의한 우측 요관-종양 샛길로 형성된 요낭종으로 진단되었고, 수술적 절제를 계획하였다. 하지만 Tc-99m diethylenetriamin pentaacetic acid 신장 신티그래피를 통하여 양측 요관-종양 샛길을 발견하였고, 치료 계획을 보존적 치료로 수정하였다.

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