Concurrent Papillary Carcinoma Arising in Thyroglossal Duct Cyst and Thyroid Gland: A Case Report1

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The occurrence and diagnosis of thyroglossal duct carcinoma is very rare. The synchronous occurrence of papillary carcinomas arising in a thyroglossal duct cyst (TGDC) and thyroid gland is extremely rare. Sistrunk’s surgical technique must always be the initial treatment for a TGDC. However, if there is an intra-thyroidal carcinoma or local invasion, thyroidectomy has to be considered. Accurate pre-operative radiological evaluation should be performed in order to plan a surgical strategy. The aim of this report was to review our experience in the management of papillary thyroid carcinoma associated with TGDC. Our patient was a 67-year-old man who had a mural, micro-calcified nodule within a palpable, thick-walled cyst at the level of the hyoid and synchronously, a small macro-calcified mass in the isthmus of the thyroid gland.

Index words: Thyroglossal cyst
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Thyroglossal duct cyst (TGDC) is the most common type of midline neck masses in children. Though usually benign, 1% of cysts may undergo neoplastic change. The most common histological subtype is papillary carcinoma (80%) followed in order of frequency by mixed follicular-papillary, squamous, follicular, anaplastic, and hurthle-cell carcinoma (1).

The synchronous occurrence of thyroglossal duct carcinoma and thyroid carcinoma in the thyroid gland is extremely rare. Debate exists as to where the original sites of primary process are located as well as whether thyroglossal duct carcinoma and thyroid carcinoma truly are synchronous. We report a case of coexistence of papillary carcinomas in the thyroglossal duct cyst and thyroid gland.

Case Report

A 67-year-old male patient presented with an anterior midline neck mass. A 10-year history was highlighted by the slowly enlarging mass in the preceding months, however no dyspnea, dysphasia, or hoarseness was noted. A physical examination revealed a 3-cm midline mass below the hyoid bone that was painless, not tender, and well-demarcated. The thyroid gland was of normal size with no palpable nodules. Further, no cervical lymph nodes were palpated. A thyroid function test revealed normal triiodothyronine, thyroxine, thyroid stimulating hormone (TSH), and thyroglobulin levels. Lastly, other laboratory investigations revealed no abnormalities.
Ultrasound (US) demonstrated a 0.8-cm mural, micro-calcified nodule within a 2.6 × 2.3 cm, partially thick-walled cyst at the level of the hyoid (Fig. 1A). In addition, there was a 1.8 × 0.8 cm, oval shaped, macro-calcified mass in the isthmus of the thyroid gland that is suggestive of a malignant nodule (Fig. 1B). A computed tomography (CT) scan demonstrated a cystic mass containing an enhancing mural nodule below the level of the hypoid bone (Figs. 1C, D). Additionally, a small dense calcified mass in the isthmus was identified adjacent to the cystic mass. On the basis of the clinical and imaging findings, our diagnosis was the coexistence of carcinomas arising in a TGDC and thyroid carcinoma.

The patient underwent a Sistrunk operation and total thyroidectomy under general anesthesia. Gross examination of the cystic mass showed a 3 cm, cystic mass filled with yellow serous fluid, and a 1.0 × 0.9 × 0.6 cm mural nodule protruded from the inner surface of the wall. The nonneoplastic cystic spaces were partially lined by flattened epithelial cells. Histopathological ex-
amination disclosed papillary carcinoma that was confined to the cyst and isthmic papillary carcinoma.

After the procedure, the patient was discharged without further significant complications. Three months after the operation, the thyroglobulin level in the blood following TSH stimulation was normal (< 1 ng/mL), and anti-thyroglobulin antibody was 1.42 U/mL (0–55 U/mL). The patient was treated with radioiodine ablation. Over the course of a 2-year follow-up from time of the procedure, ultrasound of the neck displayed no recurrence in the thyroid gland or cervical lymph nodes.

### Discussion

The median age for development of carcinoma in a thyroglossal duct cyst is 40 years, and few cases present before 14 years (2). Papillary carcinoma is the most common histologic subtype, with a higher female to male ratio (female: male = 3:2) (3). Papillary carcinoma within thyroglossal duct cyst usually presents as an asymptomatic mass. Uncommonly, dysphasia, voice change, and draining cutaneous sinuses may be developed.

The clinical features of a thyroglossal duct cyst do not usually distinguish it from a benign TGDC. Thyroglossal duct carcinoma may present with a rapidly enlarging neck mass, but the same history can be elicited in patients with recent infection in a benign cyst. Thus, in most cases a diagnosis of malignancy is not made until after surgery. Nevertheless, carcinoma should be suspected in any TGDC that is hard, fixed, irregular, or associated with lymphadenopathy (4).

The differential diagnosis of central cervical cystic lesions involves many diseases including a dermoid cyst, mucocele, thyroid diseases (benign colloid nodule, adenoma, papillary carcinoma, and cystic Hashimoto’s thyroiditis in the pyramidal lobe), a delphian lymph node with cystic thyroid papillary carcinoma, a cystic parathyroid tumor or abscess, or a branchial cleft cyst located in the midline, and a complicated TGDC (5). Some authors recommend the use of a US-guided FNAB (fine needle aspiration biopsy) in all these tumors because of the variability.

The origin of thyroid cancer in a thyroglossal duct cyst is debated. Some believe that its occurrence is de novo from islands of normal thyroid tissue found in thyroglossal duct remnants. This theory is supported by the fact that ectopic thyroid nests have been identified histologically in as many as 62% of the surgical specimens of thyroglossal duct cysts, and that carcinoma in a thyroglossal duct cyst is found without evidence of thyroid malignancy (6). Additionally, the synchronous occurrence of papillary carcinomas in a thyroglossal duct cyst and the thyroid gland can be explained as a representative of multifocal tumors. Others suggest that they are probably metastases from a thyroid carcinoma through a patent thyroglossal duct (7).

The thyroid carcinoma, concomitant with TGDC carcinoma is rare and is frequently not palpable or detectable by preoperative imaging techniques (8). A radiologic imaging study can alter the surgical plan. The US findings of TGDC may be variable, but the presence of a solid component should alert physicians to the possibility of the carcinoma. The CT findings of TGDC carcinoma are a solid nodule in the cyst, as observed our case, calcification, and an irregular margin.

The reported rate of nodal metastasis in patients with TGDC carcinoma is low. A prior study has reported that the lymphatic drainage route may differ from thyroid carcinoma (9). It is supposed that the drainage runs primarily along the superior thyroid vessels, directly to the lateral neck and without direct drainage to the central neck, because of the anatomic location of TGDC.

The standard surgical procedure for the management of thyroglossal duct anomalies is the Sistrunk procedure, which consists of excising the entire tract of the thyroglossal duct, midportion of the hyoid bone, and a portion of the base of the tongue. Thyroidectomy and regional neck dissection is indicated only if there is an intra-thyroidal carcinoma or local invasion of TGDC carcinoma.

Surgeons should be aware of concurrent TGDC and thyroid carcinomas. Moreover, a pre-operative radiologic evaluation should be performed in order to accurately plan the most effective surgical strategy. We report a rare case of the coexistence of papillary carcinomas in the TGDC and thyroid gland.

### References

감상설관낭과 감상선내에 동시에 발생한 암종: 증례 보고

김경태, 김여주, 김세중, 조영업, 전용선, 김윤정

감상설관낭 암종은 드물게 발생하지만 감상설관낭 암종과 감상샘 암종이 동시에 발생한 경우는 더욱 드물다. 감상설관낭 암종의 기본 치료는 Sistrunk 수술이지만 감상샘 내 암종이 있거나 국소 점음이 있는 낭종 내에 암종이 발생한 경우 감상선 절체가 필요하다. 이처럼 수술 전 정확한 영상의학적 진단은 수술에 큰 영향을 미친다. 이에 저자들은 감상설관낭 암종과 감상샘 유두암종이 동시에 발생한 증례를 보고하고자 한다. 67세 남자이며, 설골 부위의 만져지는 낭중 안에 벽 내 미세석화 화과 종괴와 감상샘 혈부에 작은 석회화 종괴가 함께 존재하였다.