Thyroglossal duct cyst carcinomas: is there a need for thyroidectomy?

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ABSTRACT

OBJECTIVE: Thyroglossal duct cyst (TGDC) carcinoma is a rare entity and its management is controversial. The aim of this retrospective study was to: (a) identify patients with TGDC carcinoma followed up in our clinic and (b) study specific characteristics of the disease and their association with thyroid carcinoma. DESIGN: Medical files of patients with TGDC carcinoma were reviewed and tumour characteristics, lymph node metastases, treatment and follow-up were evaluated. RESULTS: A total of 6 patients, 4 females and 2 males, mean age 39.3 years (median 33.5), were treated for papillary thyroid carcinoma arising in a TGDC. Carcinoma of the thyroid gland was found simultaneously in 4 of these patients, while in one patient thyroid carcinoma developed 10 years after the diagnosis of TGDC carcinoma. A variable clinical picture and presentation was recorded. The most aggressive manifestation of the disease in terms of local infiltration, local recurrence and lymph node metastases was observed in our youngest patients. CONCLUSIONS: Long-term follow-up is necessary for patients with thyroid carcinoma arising in a TGDC. In view of the frequent co-existence of thyroid cancer in these patients, we would recommend detailed thyroid evaluation and, eventually, total thyroidectomy at initial diagnosis of TGDC carcinoma.

Key words: Papillary carcinoma, Thyroglossal duct cyst, Thyroid cancer

INTRODUCTION

A thyroglossal duct cyst (TGDC) arises as a cystic expansion of a remnant of the thyroglossal duct tract, its incidence in post-mortem material being 7%.1 The incidence of primary carcinoma within the TGDC ranges from 0.7%2 to 1.6%3 of all patients operated on for TGDCs, although the latest published series reports a higher rate of 6.5% (9 patients with TGDC carcinomas were found among 139 patients with thyroglossal duct cysts).4 The most common tumour (90%) found in TGDCs is papillary carcinoma, followed in
decreasing frequency by squamous cell carcinoma, follicular, Hürthle cell, insular or, rarely, anaplastic thyroid carcinoma. Medullary carcinoma has not been described, as C cells are not found in the median thyroid and they are thus absent in the thyroglossal duct.\textsuperscript{3}

Simple cyst excision results in a 30-60% recurrence rate.\textsuperscript{5} Therefore, Sistrunk's procedure, consisting of excision of the cyst, the body of the hyoid bone and a core of tissue around the thyroglossal tract through the muscles of the tongue up to the foramen caecum, is recommended. With this procedure the recurrence rate is lower, in the order of 3-10%.\textsuperscript{6} Moreover, Sistrunk's procedure is considered as the minimum surgical strategy for papillary carcinoma arising in TGDCs.\textsuperscript{7,8}

Carcinoma of the thyroid gland is found in 27% to 56%\textsuperscript{7,8} of patients diagnosed with TGDC carcinoma. Two theories have been developed regarding its origin: the first theory supports the de novo occurrence of cancer in the TGDC, while the second proposes that TGDC carcinoma may represent a metastasis from a focus of cancer within the thyroid.\textsuperscript{4} As a result, there is no consensus as to whether total thyroidectomy should be performed immediately after initial diagnosis of a carcinoma arising in a TGDC. Total thyroidectomy would allow appropriate follow-up of the patients,\textsuperscript{2,9,10} although some authors do not recommend this procedure on a routine basis.\textsuperscript{7,8,11,12}

Given the rarity of the disease and the fact that the clinical picture may vary at presentation, we reviewed a series of 6 cases with TGDC carcinoma treated at our hospital. In each case, the principal features of the patient’s history are presented.

**PATIENTS - METHODS**

The medical records of 1770 patients with follicular cell derived thyroid cancer were reviewed. Among those, 6 patients with carcinoma arising in a TGDC (all of the papillary type) were identified (4 females, 2 males). The mean age of the patients was 39.3 years (median 33.5, range 18-68 years). Five out of the 6 patients also had papillary carcinoma of the thyroid: in 4 the diagnosis was made simultaneously with the TGDC carcinoma; interestingly in one patient the thyroid cancer developed 10 years after the diagnosis of TGDC carcinoma. Post-operatively, all patients received thyroxine therapy to suppress TSH and were treated with radioactive iodine (RAI). The clinical characteristics of the patients and the histological characteristics of the tumours are shown in

<table>
<thead>
<tr>
<th>Patient</th>
<th>Sex</th>
<th>Age</th>
<th>TGDC Ca size (mm)</th>
<th>Interval to thyroidectomy</th>
<th>Number of operations</th>
<th>Thyroid Ca size (mm)</th>
<th>Lymph nodes metastases</th>
<th>RAI (mCi)</th>
<th>Follow-up (years)</th>
<th>Last TG stimulated (ng/ml) **</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>18</td>
<td>8</td>
<td>10 years</td>
<td>two</td>
<td>9</td>
<td>Invasive</td>
<td>no</td>
<td>100</td>
<td>one</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>21</td>
<td>NA</td>
<td>Simultaneous removal</td>
<td>two</td>
<td>no cancer was found</td>
<td>no</td>
<td>250*</td>
<td>nine</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>18</td>
<td>NA</td>
<td>1 month</td>
<td>two</td>
<td>multifocal (1st tumor: 5, 2nd tumor: 3)</td>
<td>yes</td>
<td>250* (2 doses)</td>
<td>two</td>
<td>32.6</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>68</td>
<td>6</td>
<td>Simultaneous removal</td>
<td>one</td>
<td>6</td>
<td>no</td>
<td>100</td>
<td>three</td>
<td>0.01</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>46</td>
<td>NA</td>
<td>3 months</td>
<td>two</td>
<td>multifocal (1st tumor: 10, 2nd tumor: 3, 3rd tumor: 1)</td>
<td>no</td>
<td>100</td>
<td>seven</td>
<td>0.1</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>65</td>
<td>inv</td>
<td>6 months</td>
<td>two</td>
<td>2</td>
<td>no</td>
<td>100</td>
<td>one</td>
<td>0.6</td>
</tr>
</tbody>
</table>

TGDC: thyroglossal duct cyst, Ca: carcinoma, RAI: radioactive iodine, NA: not available
*recurrence of TGDC Ca, **on stopping thyroxine
Late occurrence of thyroid cancer in a patient with TGDC carcinoma (Patient 1)

A 28-year old male with a history of TGDC carcinoma was referred to our outpatient clinic after the diagnosis of a thyroid nodule. The patient had had surgical resection of a 3.5 cm cystic lesion in the anterior part of the neck at the age of 18 that proved to be an 8 mm TGDC carcinoma confined to the cyst wall. At that point thyroid ultrasound did not indicate abnormalities. No other treatment or follow-up had been recommended to the patient in the following 7 years. On the patient’s initiative, a neck ultrasound was performed which showed a 5 mm solid hypoechoic thyroid nodule in the left lobe; during the following 2 years this nodule doubled in size and developed ill-defined margins and microcalcifications.

At first evaluation in our clinic, a neck ultrasound confirmed the presence of an 11 mm thyroid nodule and additionally identified a 12 mm TGDC remnant. Fine needle aspiration cytology (FNAC) of the thyroid nodule was negative for cancer (category II by Bethesda).13 Due to suspicious ultrasonographic findings, the patient underwent total thyroidectomy and removal of the TGDC remnant. Histology revealed a 9 mm papillary carcinoma in the left thyroid lobe and a recurrent papillary carcinoma in the TGDC remnant (Table 1). At this point the patient received ablation with 100 mCi $^{131}$I. After 2 years of follow-up, the patient is free of local and metastatic disease, as confirmed by ultrasound, whole body scan and stimulated thyroglobulin.16

Residual disease after incomplete surgery for TGDC carcinoma (Patient 2)

A 21-year old female was referred to our department for further evaluation and follow-up after a recent diagnosis of papillary carcinoma arising in a TGDC with cyst wall invasion. The presenting symptom was a lump in the front of the neck. The patient underwent thyroidectomy at the same time, which was negative for malignancy. At this point she was referred to us for further management.

Physical examination revealed a small palpable painless mass at the level of the hyoid bone. Computed Tomography (CT) and Magnetic Resonance Imaging (MRI) scans revealed the presence of a heterogeneous, mainly solid lesion of 2.2 cm in diameter, suggestive of a TGDC remnant (Figure 1 a-f). Re-operation was recommended and histology identified...
an invasive 5 mm papillary carcinoma in the TGDC remnant (Figure 2 a, b). Ablation with 100 mCi of $^{131}$I was given, followed by a second dose 9 months later due to residual local disease (Table 1). During 10 years of follow-up, no recurrences or metastases were identified. She is now on replacement therapy with thyroxine therapy and has recently given birth to her first child.

Simultaneous diagnosis of TGDC and thyroid gland microcarcinomas (Patient 4)

A 68-year old female was referred to our department for evaluation and further treatment. The patient had a 10-year history of multinodular goiter, which had recently been complicated by hyperthyroidism. Additionally, she was followed up for a midline cystic neck mass that had progressively increased in size and was now 3 cm in its maximal diameter. Ultrasound of the thyroid confirmed the existence of the goiter and the presence of small isoechoic nodules as well as the cystic mass in front of the hyoid bone. Excision of the cystic mass was decided and total thyroidectomy of the hyperfunctioning multinodular gland was performed in parallel. Histology revealed a papillary carcinoma arising in a TGDC of 6 mm within the cyst wall and the thyroid gland also harboured one focus of papillary 6 mm microcarcinoma (Table 1).

The need for thyroid evaluation before operation for TGDC (Patient 5)

A 46 year-old female was referred to our department 4 months after the diagnosis of TGDC carcinoma confined to the cyst wall. On clinical examination, the thyroid gland was enlarged and neck ultrasound revealed a multinodular goiter. FNAC of a 12 mm
thyroid nodule was suspicious for papillary carcinoma (category V by Bethesda). The patient underwent total thyroidectomy and histology showed multifocal papillary carcinoma (3 tumour loci of 10.3 and 1 mm). After surgery the patient received an ablative dose of 100 mCi of radioactive iodine. His biggest complaint was that the two operations were not performed at the same time.

**Preoperative diagnosis of TGDC carcinoma with FNAC (Patient 6)**

A 65-year old male with a recent diagnosis of papillary carcinoma arising in a TGDC, invasive to surrounding fat tissue, was referred to our department for further evaluation and treatment (Figure 3). Preoperatively, the patient had a long-standing growing midline mass, which on CT scans before and after IV contrast was solid with homogeneous enhancement (Figure 4 a, b). FNAC of the cyst was suspicious for malignancy (category V by Bethesda system).

Thyroid ultrasound at presentation showed hypoechoic nodules in both lobes. FNAC of the dominant nodule was negative for malignancy (category II by Bethesda). However, due to the need for RAI therapy of the invasive TGDC carcinoma the patient was referred for total thyroidectomy. Histology revealed a 2mm papillary microcarcinoma in the isthmus (Table 1). Subsequently he received ablation with 100 mCi of radioiodine.

**DISCUSSION**

In the present study, we report a 66% co-existence of thyroid cancer after initial diagnosis of TGDC carcinoma. The prevalence of co-existing malignancies in both TGDC and the thyroid increased to 83% after longer follow-up, as one patient was diagnosed with thyroid cancer 10 years after the TGDC carcinoma diagnosis, a rate that is higher than that reported in other studies.\(^7\)\(^9\) This could be attributed to selection criteria and long-term follow-up, as in 5 of our patients thyroid surgery was decided due to abnormalities in the thyroid gland. Patel et al\(^8\) evaluated 5 patients with TGDC carcinoma, this being in addition to the 57 cases published in the literature, and found thyroid carcinoma in 27%. In these cases, thyroid surgery was performed as part of the initial treatment of TGDC.
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rounding soft tissue invasion, lymph node metastases and prognosis), most authors recommend that the management of TgDC carcinoma should be based on current guidelines, as for any other differentiated thyroid cancer.7,12,17

One of our patients (case 1) was diagnosed with thyroid carcinoma 10 years after the initial diagnosis of TgDC carcinoma. It is not certain whether a focus of thyroid carcinoma existed at the time of diagnosis of TgDC carcinoma or whether it developed at a later time. Based on ultrasound findings, we believe that the thyroid carcinoma developed later on. The development of thyroid cancer 10 years after the diagnosis of TgDC carcinoma has rarely been reported in the literature.9 Three of our patients were younger than 21 years of age, this emphasizing the importance of adequate screening and long-term follow-up in young patients who present with TgDCs. In this series, an increased female to male ratio is seen, as in cases of differentiated thyroid cancer.

The long-standing controversy as to whether thyroid cancer arising in a TgDC represents a metastasis from thyroid cancer or not has not as yet been resolved with certainty, especially in cases of simultaneous discovery of both cancers. In other cases, as in our first patient, the possibility that carcinoma in TgDC is a metastasis is small, as thyroid examination after the TgDC excision did not reveal thyroid morphological abnormalities. Thyroglossal ducts contain, by definition, thyroid epithelium and thus may develop primary cancer. Similarly, it is difficult to identify whether in cases of multifocal thyroid cancer: a) there is a primary thyroid lesion which causes intrathyroidal metastases or b) if all thyroid tumour sites are developed simultaneously and independently from each other.

An interesting point that should be discussed is that, similar to the published series, in our patients the thyroid malignancies were all microcarcinomas.18 A plausible explanation for this finding is the fact that during close follow-up after the TgDC carcinoma diagnosis, thyroid cancer is detected and excised earlier. In our series, despite the presence of small size of thyroid tumours, invasive and multifocal thyroid carcinomas were also observed.

One possible weakness of our study is the fact that
the number of patients is relatively small. However, 
TGDC carcinoma is a rare entity, with around 250 
cases having been published in the world literature. 
Thus, most published studies are small retrospective 
case series like our own. Even the most recent series 
emphasize the weak results of research due to the 
small numbers of patients. 19

Based on the available data, we could conclude that 
ultrasound of the thyroid gland must be performed 
in all patients. If nodule(s) are identified, further 
evaluation is recommended. In the event of suspi-
cious ultrasonographic or cytological findings, total 
thyroidectomy must be applied. Otherwise, long-term 
follow-up is necessary to exclude the development of 
thyroid cancer. Further treatment of both cancers 
must be based on current guidelines for differenti-
ated thyroid cancer.

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