ABSTRACT

Warthin-like tumor of the thyroid is a recently described rare variant of thyroid papillary carcinoma. The distinguishing histological feature of this variant is papillary foldings lined by oncocytic neoplastic cells with clear nuclei and nuclear pseudoinclusions, accompanied by prominent lymphocytic infiltrate in the papillary stalks. Its prognosis has been reported to be almost similar to conventional papillary carcinoma. In this case series, we report four cases with Warthin-like papillary carcinoma of the thyroid, diagnosed at Dokuz Eylül University Faculty of Medicine Department of Pathology in 2008 and 2009. Three patients were female. The mean patient age was 39 years (range, 20-56) and the mean tumor size was 1.7 cm (range, 0.9-2.0 cm). All of the cases had lymphocytic thyroiditis in the background. None of the tumors showed lymphovascular invasion. The patients are free of any recurrence and/or distant metastasis with a mean follow-up of 25 months. This rare variant of thyroid papillary carcinoma with distinct histopathological features should be indicated in pathology reports. Further studies and long-term follow-up of patients are needed to highlight the biological behavior of this variant.

Key Words: Warthin tumor, Papillary carcinoma, Thyroid, Lymphocytic thyroiditis, Oncocytic change

INTRODUCTION

Papillary carcinoma is the most common primary malignant neoplasm of the thyroid gland. According to the current classification of thyroid tumors by the World Health Organization (WHO), the major morphologic variants of papillary carcinoma are follicular, macrofollicular, oncocytic, clear cell, tall cell, columnar cell, diffuse sclerosing and solid variants (1). Warthin-like variant of papillary carcinoma is an uncommon variant that was first described by Apel et al. in 1995 (2). This morphologic variant has distinct features of papillary architecture with prominent lymphocytic stroma in fibrovascular cores, resembling Warthin tumor of the salivary gland. The neoplastic cells lining the papillary folds have nuclear features of usual papillary carcinoma. Studies show that the overall and progression-free survival rates of these patients are similar to patients who have conventional papillary carcinoma (2-17). There is only one case of Warthin-like tumor of papillary carcinoma reported in the literature, who underwent anaplastic changes (14).

In this paper, we report four cases of Warthin-like tumor of papillary carcinoma, diagnosed at our institution in 2008 and 2009. The clinicopathological features of these patients are discussed, and the previously published reports are reviewed.
CASE REPORTS

Four cases diagnosed at Dokuz Eylul University Faculty of Medicine Department of Pathology in 2008 and 2009 were selected from the pathology archives of our institution. The clinical and follow-up information of these patients, including age, sex, size and localization of the tumor, and extent of surgery were obtained from the medical charts of the patients. Fine-needle aspiration (FNA) cytology was performed in three cases. The mean follow-up period of the patients was 25 months (range, 22-26 months).

The clinical features and the histopathological findings of the patients are summarized in Table I and Table II, respectively. The mean patient age was 39 years (range, 20-56). Three cases were female and one case was male. FNA cytology of two cases were diagnosed as suspicious for malignancy, and the other case as malignant (Figure 1). All of the cases underwent total thyroidectomy. Lymph node biopsy was performed in two cases (cases 1 and 3), in whom metastasis was not identified. Radionuclide therapy was performed in all patients. Postoperatively, each patient was clinically followed for local recurrence and/or distant metastases both by radiologic and scintigraphic methods. The mean follow-up period of the patients was 25 months (range, 22-26 months), and there was no sign of either local recurrence and/or distant metastases in any of the patients.

Multifocality was detected in three cases. The largest tumor size was 2 cm in three cases and 0.9 cm in one. One case showed invasion of the thyroid capsule and peripheral soft tissue. None of the tumors showed lymphovascular invasion at the time of diagnosis. Histologically, all of the tumors showed papillary architecture with lymphoplasmocytoid stroma in the papillary stalks (Figure 2). The neoplastic cells lining the papillary fronds had clear nuclei with a few nuclear pseudoinclusions, and abundant granular eosinophilic cytoplasm (Figure 3). All of the cases had lymphocytic thyroiditis in non-neoplastic areas (Figure 4).

DISCUSSION

Warthin-like papillary carcinoma is an uncommon variant of thyroid papillary carcinoma. Major histological features of these tumors are neoplastic cells with oncocytic changes and clear nuclei, lining the papillary fronds that have lymphocytic stroma in their stalks. The name comes from its close resemblance to Warthin tumor of the salivary gland. When Apel et al. first described the features of these tumors in 1995, they named it as “papillary Hurthle cell carcinoma of the thyroid with lymphocytic stroma - Warthin-like tumor of the thyroid” (2). After their description, approximately 95 cases have been reported in the literature, mostly as single case reports (2-16).

The mean age of patients with conventional papillary carcinoma is the mid 40’s to early 50’s (1). However, Warthin-like tumor variant of papillary carcinoma have been reported in a wide age range (19-85 years) in the literature (2-16). The mean age of our cases was 39 years (range, 20-56 years). Although some authors claim that this variant is mostly seen in elder patients, we believe it is more reasonable to expect these tumors at a broader age group than the conventional papillary carcinoma (11). Female/
male ratio is high in these tumors similar to the other papillary carcinoma variants. We had only one male patient as opposed to three females.

Previous case series of Warthin-like tumor variant of papillary carcinoma, together with the cases presented here, are summarized in Table III. It is emphasized that these tumors behave like usual papillary carcinomas with excellent long term overall survival (2-17). To the best of our knowledge, the only case of Warthin-like tumor variant of papillary carcinoma with anaplastic changes reported in the literature is a 74 year-old female (14). Having a recurrent nerve and lymph node involvement at the time of diagnosis, by a delay of three years in her treatment, she died of the disease 18 months after the operation. One of our cases showed extrathyroidal extension, that is accepted as an unfavorable prognostic factor by the current WHO classification (1). With a mean follow-up period of 25 months, none of our cases showed any local recurrence and/or distant metastasis. However, this time period is limited to make a realistic comment about the clinical behavior of these tumors.

The histologic appearance of Warthin-like tumor variant is similar to tall cell and columnar cell variants of papillary carcinoma that tend to show more aggressive clinical

Figure 1: FNA cytology of case 2 showing pleomorphic neoplastic cells with eosinophilic cytoplasm, one of which has a pseudonuclear inclusion (H-E, x400).

Figure 2: Histological appearance of case 4. There is prominent lymphocytic infiltrate in the stalks of the papillary folds (H-E, x100).

Figure 3: High power view of the same case as in Figure 2 showing neoplastic oncocytic cells with clear nuclei and a few nuclear grooves lining the papillary folds (H-E, x400).

Figure 4: Lymphocytic thyroiditis in non-neoplastic areas of case 1, accompanying the tumor (H-E, x100).
behavior than usual papillary carcinoma (18-23). The height of the oncocytic tumoral cells in the tall cell variant of papillary carcinoma is at least three times their width (1). The columnar cell variant, on the other hand, is composed of columnar cells containing cytoplasmic vacuoles, but they lack lymphocytic stroma (23). Various papillary carcinoma variants as well as some other histologic types of thyroid carcinoma, including medullary carcinoma may show oncocytic changes (24,25). Besides, Hurthle cell carcinoma of the thyroid, which has worse prognosis than papillary carcinoma and its variants, may have pseudopapillary architecture histologically, although the nuclei lack usual papillary carcinoma features (26). The distinguishing feature of Warthin-like tumor of papillary carcinoma from all other histologic types and subtypes of thyroid carcinoma is the existence of lymphocytic stroma in the papillary stalks. This stroma may also contain follicles with germinal centers (12).

Some authors consider existence of lymphocytic stroma in a thyroid tumor as an indicator of favourable prognosis (9). According to these authors, tumor associated lymphocytes may play a role as a control mechanism. However, this claim has not been proven yet. Besides, lymphocytic infiltrate is commonly seen in thyroids with or without neoplasm (27). The case reports in the literature point out that lymphocytic thyroiditis or Hashimoto thyroiditis is an associating condition in most of the Warthin-like tumors (2-17). The role of intratumoral lymphocytes, whether as an autoimmune response impeding the spread of the tumor or providing a generous background for the development of the neoplasm, remains controversial (29-39).

FNA cytology findings of these tumors were first described by Yousef et al. (3) and Baloch et al. (40). The differential diagnosis of oncocytic variant of papillary carcinoma, conventional papillary carcinoma, Hurthle cell neoplasm arising in a background of lymphocytic thyroiditis, and Warthin-like papillary carcinoma can be challenging due to common findings in FNA. The pleomorphic oncocytic cells with nuclear features reminiscent of papillary carcinoma, admixed with inflammatory cells in the background should be warning in terms of Warthin-like papillary carcinoma (3,5,40). Among our three cases who underwent FNA, cytological features supported malignancy in one, and were suspicious for malignancy in two cases. In evaluating FNA cytology, the most important thing is not to consider the prominent inflammatory cells in the background as “lymphocytic thyroiditis”, which may give rise to false negative diagnosis.

Due to the heterogeneity of thyroid papillary carcinomas, areas with features of usual or rare histologic variants may be seen in the same tumor. There are studies that report coexistent Warthin-like tumor variant of papillary carcinoma and other variants of thyroid carcinoma in the same patient. Some of these reported cases had worse prognosis due to the unfavourable prognosis of the accompanying tumors (26). In all of our cases, there was either a conventional, oncocytic or follicular variant of papillary carcinoma accompanying the tumor.

Multifocality is a frequent finding in papillary carcinomas and does not have any prognostic implication (1). Three of our cases were multifocal. One of the cases with multifocal tumor also had thyroid capsule and peripheral soft tissue invasion. Thyroid capsule invasion seems to be an unfavourable prognostic factor (1). However, it is not well known whether multifocality is a risk factor for the tumor to invade the thyroid capsule.

In conclusion, papillary carcinomas usually have perfect clinical outcome. Nevertheless, identifying the prognostically different subtypes of papillary carcinoma with distinct histopathological features is significant. We
believe that Warthin-like tumor variant of papillary thyroid carcinoma, as well as other variants, should be indicated in pathology reports. Further studies and long-term follow-up of the patients are needed to highlight the biological and clinical behavior of these tumors.

REFERENCES


