Columnar Cell Variant of Papillary Thyroid Carcinoma: A Clinicopathologic Analysis

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Abstract: Background: Papillary Thyroid Carcinoma (PTC) is the Most common thyroid malignancy disease, and it is more common in females. It happened in any age. Mean age at diagnosis is approximately 40 years old. There is history of neck irradiation in 5–10% patients. And non-neoplastic gland may show nuclear aberrations as a result. There is increased incidence in Hashimoto's thyroiditis, but not clear whether there is an increased incidence in Graves' disease. Columnar cell variant of papillary thyroid carcinoma is the rare subtype of PTC, which is reportedly about 0.17%. CCV-PTC was reported in 1986 by Evans for the first time. It is different from the typical papillary carcinoma, not only in morphology but also in biological behavior, and is more aggressive than typical papillary carcinoma. Objective: To study the clinical-pathological links and features of papillary thyroid carcinoma (Columnar cell variant) (CCV-PTC). Methods: 4 cases of CCV-PTC from Chengde medical college affiliated hospital were retrospectively analyzed, to observe its clinical features, histological, immunophenotype, and metastasis. And the literature of CCV-PTC were reviewed. Results: Out of the 4 patients, two male patients of 70 and 53 years old, two female patients of 64 and 68 years old. Microscopically, tumor cells arranged in multistage papillary structure, with pseudostratified columnar epithelium, same to gastrointestinal and lung metastatic carcinoma. IHC: Ki-67(5-15%), calcitonin(-), TG(+), ck19(-), TTF-1(+), CgA(-), SyN(-), ck(+/-). Conclusion: CCV-PTC is a special subtype of PTC with a more aggressive biological behavior, particularly among older people, Immunohistochemical method is helpful for the diagnosis.

Keywords: Papillary Thyroid Carcinoma, Columnar Cell Variant, Pathological Feature

1. Introduction

Papillary thyroid carcinoma is the most common thyroid cancer, about 60%-70%, and often encountered in children and adolescents, especially young women [1-3]. Age has a significant impact on prognosis, which is good in young patients and poor in old patients. The histologic subtypes of papillary thyroid carcinoma included follicular variant, diffuse sclerosing variant, columnar cell variant, tall cell variant, oxyphilic cell variant, Warthin’s-like variant, papillary thyroid carcinoma with nodular fasciitis-like stroma., cribriform papillary thyroid carcinoma, and radiation-induced pediatric thyroid cancer. CCV-PTC is a rare type of papillary thyroid carcinoma of high malignancy. 4 cases of CCV-PTC were collected, to discuss the histopathological characteristics, genetics, and prognosis.

2. Materials and Methods

2.1. Clinical Data

Case 1 Female, 64 years old, the tumor was located in the right leaf of the thyroid, about 3.0cm x 3.0cm. Boundary clear, smooth surface, without tenderness were the mainly clinical symptoms. B-ultrasonography showed several hypoechoic nodules, weak echo nodular and hybrid echo-mass in thyroid parenchyma of double sides. The largest nodule was
25.3mm×16.8mm×24.1mm in right side, with clear boundary and relatively regular shape. CDFI: color flow was found surrounding the nodule.

Case 2 Male, 70 years old, with III degree swelling of thyroid. Multiple nodules was in left side, out of which the largest one was 8.0cm×15.0cm. Boundary clear, smooth surface, without tenderness were the mainly clinical symptoms. No noise of blood vessels was heard. Palpable cervical lymph node, proptosis and hands shaking were not found. CT showed slightly high density shade under sternocleidomastoid muscle of the left, about 5.5x6.3x10cm, heterogeneous enhancement and punctate calcifications.

Case 3 Male, 58 years old, the tumor was located in the right leaf of the thyroid, about 6.5cm×5.5cm. Boundary clear, smooth surface, without tenderness were the mainly clinical symptoms. B-ultrasonography showed several medially echoic-mass in thyroid parenchyma of left sides, with clear boundary and relatively regular shape. Medially echoic-mass, about 60.0mm×28.5mm×63.6mm, was found in supravacuicular fossa.

Case 4 Female, 68 years old, 10 cm neoplasm was found in the neck, with unclearly boundary and very tender. CT showed high density shade in the soft tissue of front clavicle.

2.2. Methods

4 cases of specimens are fixed in the neutral formalin liquid for 12 hours. The samples were taken for routine histopathological observation. IHC was used to be diagnosed definitively. Ki-67, calcitonin, TG, ck19, TTF-1, CgA, SyN and CK were from Jinqiao Company in Beijing, and compared with the positive control group and negative control group.

3. Results

3.1. Macroscopic

Case 1 Broken tissue, about 1.5x1x0.5cm, dusty red, Calcifications was seen in part area. Case 2 A gray-red nodular mass, about 10x5x4cm, with intact capsule. Case 3 Nodule, about 6.5×5×5cm, with clear boundary, gray-red, firm. Case 4 Nodule, about 13x7x3.5cm. gray-red.

3.2. Microscopically

Tumor cells arranged in multistage papillary structure, with pseudostratified columnar epithelium, same to gastrointestinal and lung metastatic carcinoma. (Figure 1-4).

3.3. Immunohistochemistry

TG and TTF-1 were positive in tumor cells of CCV-PTC, while Calcitonin, CK19, CgA, SyN were negative. CK was weakly positive. Ki - 67 (5-15%). (Figure 5-10).

3.4. Genetics

Gene rearrangements of the PTC was RET/PTC, include RET/PTC1, RET/PTC2 and RET/PTC3, which were associated with the histological subtypes, such as RET/PTC1 in occult thyroid cancer and PTC, RET/PTC3 in tall cell variant. Mutated BRAF was found in CCV-PTC [4].

Figure 1. Tumor cells arranged in multistage papillary structure.

Figure 2. Invasive tumor cells HE 200X.

Figure 3. Nuclear vacuolization in tumor cells.
4. Discussion

Papillary thyroid carcinoma (PTC) is the most common malignancy in the thyroid gland, which accounts for 90% of all thyroid cancers. The variant of papillary carcinoma of the thyroid represents an unusual neoplasm whose clinicopathological features and biological behavior have not been thoroughly characterized. Columnar cell variant of papillary thyroid carcinoma is the rare subtype of PTC, which is reportedly about 0.17% [5]. The revised American Thyroid Association guidelines recently categorized the PTC variants according to their biological behavior as described in the literature, and CCV-PTC was classified as the aggressive type.
CCV-PTC was reported in 1986 by Evans for the first time [6]. It is different from the typical papillary carcinoma, not only in morphology but also in biological behavior, and is more aggressive than typical papillary carcinoma.

4.1. Clinical Symptoms

CCV-PTC is seen in older women, who with a worse prognosis than young women [7]. The Surveillance of 765 cases of CCV-PTC was from 1988 to 2013 by C Jiang [8]. The study showed, Compared with PTC, CCV tumors tended to be larger, with a higher incidence rate among males and in patients ≥65 years of age. CCV was associated with higher rates of extrathyroidal extension, multifocality, lymph node examinations, and lymph node and distant metastases (p<0.0001). Significant differences were found in 10-year overall survival (97.14% vs 89.15%, p<0.0001) and disease-specific survival (99.08% vs 93.07%, p<0.0001) between PTC and CCV. Clinical Differential Diagnosis 1. Thyroid Adenoma Thyroid adenoma is often seen in young women. Boundary clear, smooth surface, without tenderness are the mainly clinical symptoms. The neck ultrasound is helpful to establish the diagnosis. 2. Nodular Goiter Multiple nodules or solitary nodule in thyroid tissue. 3. Thyroid Cancer tends to have more advanced tumors than others with swollen lymph glands, hoarse voice and difficulty swallowing.

4.2. Macroscopic

There is a certain relationship between macroscopic and aggressive, such as tumor size, capsule and tumor boundary. According to the macroscopic, CCV-PTC is divided Into inert columnar cells papillary carcinoma (tumor diameter smaller, no capsular invasion, boundary clear). Conversely, if the tumor diameter, no envelope is more aggressive.

4.3. Microscopically

Among the aggressive variants of PTC, CCV-PTC is the most misdiagnosed and underrecognized entity. On histology, it is defined by papillae or gland-like structures lined by columnar cells displaying prominent nuclear stratification.

Tumor cells arranged in multistage papillary structure, with pseudostriatified columnar epithelium, round nucleus, cell hyperplasia and cytoplasmic spaces, same to gastrointestinal and lung metastatic carcinoma. Spindle cells and follicle suggested poor prognosis. It was difficult to different CCV-PTC from tall cell variant. It was also arranged in pseudostratified columnar epithelium in tall cell variant, but acidophilic granular cytoplasm, and nucleus on the basal part of cells.

4.4. The Role of Immunohistochemical in the Differential Diagnosis of CCV – PTC

TG and TTF-1 were positive in tumor cells of CCV-PTC, which negative in gastrointestinal and lung metastatic carcinoma. CEA, EMA and CK19 were negative in tumor cells of CCV-PTC,which can different from tall cell variant(EMA,CK19,CD115 all positive).

4.5. Genetics

The mutation rate of BRAF gene, located in V600E, was high in CCV-PTC, about 33%[2].There were 3 cases of CCV-PTC found BRAF mutation out of 9 cases, 2 cases with clinical invasive. IHC showed that, the expression of cyclin D1 and Ki – 67, in the nucleus, were higher, which P53 weakly positive in in invasive and inert CCV – PTC. The expression of β-catenin and Bcl – 2 were low. The expression of ER, PR in both invasive and inert CCV-PTC were high, and had nothing to do with age, gender. The expression of CDX2 was about 55%, while low expression in other types of thyroid cancer. [9, 10].

4.6. Survival Analysis

A study [5] including 48 patients of CCV-PTC showed that, 20 cases were clinically indolent, and in 23 cases the tumors were considered as aggressive. Of the cases with clinical follow-up, almost all the patients with indolent tumors (18 out of 19) were alive or free from the disease for 9 months to 22 years after diagnosis. Of the 20 patients with aggressive tumors, 13 died from disease approximately 7–126 months after diagnosis. Also, the extrathyroidal extension of clinically indolent CCV-PTC were not reported while most of clinically aggressive CCV-PTC showed extrathyroidal extension, accounting for 67–100% [11]. In the study of Cho, JH Shin [7], six cases of CCV-PTC were identified, the four indolent tumors in study were small (mean size: 1.2 cm), encapsulated, confined to the thyroid gland, and present in younger individuals (range: 27–34 years). Recurrence was not observed in the female patients during the clinical follow-up. Most individuals in the indolent group had nodules with a smooth margin based on US. On the other hand, two patients had aggressive tumors that were larger (1.8 cm and 6.0 cm) than the indolent tumors, extrathyroidal extension, metastasis to LNs and distant organs, affected older patients (55 years and 70 years) and died of their disease, 4 years after diagnosis.

5. Conclusion

The prognosis of papillary thyroid carcinoma is good. 10 – year survival rate more is than 90%, especially in young patients (98%) [12-17]. The factors affecting the prognosis included vascular invasion, nuclear atypia, tumor size and age. The prognosis of CCV-PTC is worse, because of the local invasion, lymph node metastasis and distant metastasis [18, 19]. It should be highly valued on the realization and diagnosis of CCV – PTC, to guide clinical treatment.

References


