Ultrasonographic imaging of papillary thyroid carcinoma variants

Jung Hee Shin

Department of Radiology and Center for Imaging Science, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Korea

Ultrasonography (US) is routinely used to evaluate thyroid nodules. The US features of papillary thyroid carcinoma (PTC), the most common thyroid malignancy, include hypoechogenicity, spiculated/microlobulated margins, microcalcifications, and a nonparallel orientation. However, many PTC variants have been identified, some of which differ from the classic type of PTC in terms of biological behavior and clinical outcomes. This review describes the US features and clinical implications of the variants of PTC. With the introduction of active surveillance replacing immediate biopsy or surgical treatment of indolent, small PTCs, an understanding of the US characteristics of PTC variants will facilitate the individualized management of patients with PTC.

Keywords: Ultrasonography; Thyroid cancer, papillary; Papillary thyroid carcinoma variant; Thyroid neoplasms

Introduction

Papillary thyroid carcinoma (PTC) has an excellent prognosis, with an overall cancer-related 10-year survival rate of nearly 90% [1,2]. Many variants of PTC have been documented [3]. With the existence of nuclear features, each variant has a combination of specific growth patterns, cell types, and stromal changes. Some variants are more aggressive, whereas others are more indolent, which contributes to risk stratification. Recently, the revised American Thyroid Association guidelines classified PTC variants according to their biological behavior as described in the literature [4] (Table 1). The histopathological variants of PTC associated with more unfavorable outcomes are the tall cell variant (TCV), the columnar cell variant, and the hobnail variant. Although the data remain controversial, the solid variant and the diffuse sclerosing variant (DSV) may be associated with less favorable outcomes. In contrast, the encapsulated follicular variant (FV), the cribriform-morular variant (CMV) (which is related to familial tumor syndromes), and the recently described Warthin-like variant (WV) are associated with favorable outcomes. However, the guidelines contain no information about the imaging characteristics of the variants of PTC.

Via high-resolution ultrasonography (US), thyroid nodules can be detected and, additionally, some malignant histological types can be specifically diagnosed [4–7]. Since US is the primary means of assessing thyroid nodules and provides a basis for surgical decisions, the US imaging features of the variants of PTC and their clinical implications will be described herein.
Tall Cell Variant
The TCV is composed of more than 50% tall cells, which are twice or three times as tall as wide and have eosinophilic cytoplasm and characteristic PTC nuclear features [8]. Some pathologists recommend noting the percentage of tall cells in the pathology report when it is significant (≥30%), irrespective of the cutoff used to designate a tumor as a TCV. The incidence of TCV has been reported to range from 4% to 17% of PTCs [9]. The TCV can be missed by pathologists not only because of its rarity but also due to its blurred definition. The TCV seems to be more aggressive even when it lacks extrathyroidal extension. TCV without extrathyroidal extension has been found to show higher rates of recurrence and distant metastasis than classic PTC without extrathyroidal extension (14.9% vs. 6.5% and 6.4% vs. 0%, respectively) [9]. The poorer prognosis of the TCV reflects the older age of patients at presentation, the larger size of the tumor, and a higher frequency of extrathyroidal extension [10]. The BRAF V600E mutation is found in up to 92.6% of TCV tumors [11].

On ultrasonography, TCV tumors often appear as microlobulated, markedly hypoechoic nodules with microcalcifications and extrathyroidal extension, and are always associated with lymph node metastasis (Fig. 1). TCV PTC should be included in the differential diagnosis of aggressive thyroid tumors because it almost always appears as a highly suspicious nodule on US [12].

Columnar Cell Variant
In earlier publications, the columnar cell variant was thought to always be aggressive; however, the encapsulated form is now associated with a favorable outcome after complete excision. The poor prognosis and aggressive behavior of the columnar cell variant warrant its early recognition.

The columnar cell variant is lined by elongated hyperchromatic pseudostratified columnar cells. It is very rare, with only approximately 50 cases reported since its first description in 1986 by Evans [13]. Its prognosis is variable: more favorable when the tumor is small, encapsulated, and circumscribed, and less favorable when the tumor is large and metastatic with extrathyroidal extension [14]. The BRAF V600E mutation is found in one-third of these tumors [15].

Little information has been published about the imaging features of columnar cell variants. On US, encapsulated tumors appear as circumscribed hypoechoic nodules with or without microcalcifications (Fig. 2), and aggressive tumors as large microlobulated hypoechoic nodules, often with capsular protrusions representing extrathyroidal extension and neck nodal metastasis.

Hobnail Variant
The hobnail variant has apically placed nuclei and a surface bulge in >30% of its cells [16]. This rare variant is more closely associated

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**Table 1. Classification of papillary thyroid carcinoma variants according to biological behavior in the 2015 American Thyroid Association guidelines**

| Aggressive | Less favorable | Favorable |
|------------|---------------|-----------|
| Tall cell variant | Solid variant | Follicular (encapsulated) variant |
| Columnar cell variant | Diffuse sclerosing variant | Cribriform-morular variant |
| Hobnail variant | | Warthin-like variant |

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**Fig. 1. Tall cell variant of papillary thyroid carcinoma in a 19-year-old female woman.**

A. Longitudinal ultrasonography shows a microlobulated hypoechoic nodule (arrows) with microcalcifications and extrathyroidal extension in the right thyroid gland. B. Metastatic lymph nodes (arrows) are coexistent in the lateral neck.
with recurrence, metastasis, and cancer-related death than is classic PTC [17].

In a previous report, eight of 10 hobnail variant cases had the *BRAF V600E* mutation, whereas none had an *ALK* fusion or *TERT* promoter mutation. In a literature review of 55 hobnail variant cases, most patients presented with advanced-stage cancer, and the disease-specific survival rates were 83%, 71%, and 54% at 5, 10, and 20 years after the initial operation, respectively [18]. The imaging features of the hobnail variant have not been reported owing to its rarity. This variant has been found to show a palpable mass appearing as a microlobulated hypoechoic nodule with microcalcifications and multiple metastatic lymph nodes on US.

**Less Favorable Variants**

**Solid Variant**
The solid variant comprises approximately 3% of all PTCs [19], and there are conflicting reports about its behavior in the literature. It has been identified in more than 30% of the survivors of the Chernobyl nuclear accident [20,21]. Compared with well-differentiated tumors, the solid variant appears to have a slightly greater rate of distant metastasis, a less favorable prognosis, and a less favorable survival rate. Among children and adolescents with post-Chernobyl PTCs, the percentage of solid variants was very low (<1%) during the first 10 years of follow-up [21]. In a recently published report, none of the patients with solid-variant PTCs had a history of radiation exposure [22]. The authors concluded that the clinical characteristics of the solid variant, including the frequency of lymph node metastasis, distant metastasis, and recurrence, did not differ from those of classic PTC, presumably due to the small size of the tumors (all tumors in their series were <2.0 cm) and the early detection of the thyroid cancer. Early detection of small PTCs via imaging and preoperative fine-needle aspiration appears to be important.

Histologically, more than 50% of solid variants have a solid growth pattern, with nests of non-necrotic tumor cells surrounded by a thin rim of capillary-rich connective tissue and the intact nuclear features of classic PTCs [19]. The solid variant should be distinguished from poorly differentiated thyroid carcinomas, which have similar insular, solid, and trabecular growth patterns and a lower survival rate [20]. These tumors have been diagnosed as high- or intermediate-suspicion nodules using US based on the Korean Thyroid Image Reporting and Data System [23,24]. Some tumors have been reported to have a well-defined border and could be predicted to exhibit indolent behavior [25].

**Diffuse Sclerosing Variant**
As described by the World Health Organization, DSV PTC tends to occur in young women and is characterized by the diffuse involvement of one or both thyroid lobes, usually without formation of a dominant mass, and by extensive squamous metaplasia, numerous psammoma bodies, dense lymphocytic infiltration, and stromal fibrosis.

Fig. 2. Columnar cell variant of papillary thyroid carcinoma in a 27-year-old woman. A, B. Transverse (A) and longitudinal (B) ultrasonography shows a circumscribed hypoechoic nodule (arrows) with calcifications and cystic changes in the left thyroid gland, suggesting an encapsulated tumor.
The DSV was proposed as a new PTC variant by Vickery et al. in 1985 [26] and officially recognized as a histological variant in 1988 [27].

The prognostic implications of the DSV remain controversial. Some studies reported a high incidence of lymph node metastasis and lung metastasis at presentation, thus suggesting that the DSV has a less favorable prognosis than does classic PTC [28–30]. Other reports, however, found no differences [31,32].

A recent meta-analysis confirmed the aggressive biological behavior of the DSV. When preoperatively suspected, DSV PTC should be treated via total thyroidectomy with lymph node excision followed by radioiodine therapy [33]. In a large multicenter study, the DSV had a 7-year recurrence-free survival rate of 63%, which is significantly worse than that of other PTC variants. Despite aggressive surgical interventions, the risk of recurrence of DSV PTC requires constant meticulous surveillance, similar to the follow-up protocol for other high-risk PTC patients, during the first 5 years after treatment [34].

RET/PTC rearrangement is the major genetic alteration in DSV PTC, whereas the prevalence of the BRAF mutation is relatively low (24%) [35]. The US features of the DSV include a heterogeneous echotexture in the parenchyma, an ill-defined solid hypoechoic nodule when formed as a mass, scattered microcalcifications within or without the nodule (appearing as a "snowstorm"), and representative psammoma bodies on pathology (Fig. 3) [36]. Lateral nodal metastasis with microcalcifications is frequently detected.

It is sometimes difficult to differentiate DSV PTC from chronic thyroiditis manifesting with only microcalcifications. DSV PTC is common in young patients and is always accompanied by cervical metastatic lymph nodes, whereas chronic thyroiditis is common in middle-aged patients and frequently accompanied by benign reactive lymph nodes. Patients with chronic thyroiditis have a history of hypothyroidism.

**Favorable Variants**

**Follicular Variant**

The FV is the most common type of PTC after classic PTC [37]. It possesses both the nuclear features of PTC (e.g., nuclear clearing, grooves, and pseudo-inclusions) and a follicular growth pattern [38]. FV tumors are divided into two main subgroups—infiltrative and encapsulated—that differ in their molecular profiles and biological properties. Infiltrative FV PTCs frequently harbor BRAF mutations and exhibit aggressive behavior, whereas encapsulated FV PTCs have a high frequency of RAS mutations and indolent behavior [39]. Infiltrative FV PTCs do not differ significantly from conventional PTCs in terms of prognostic factors or survival rate.

Some investigators have suggested that US can predict the behavior of the FV because FV PTC has dichotomous US features (PTC-like vs. follicular neoplasm-like). FV tumors with malignant US features seem to behave more aggressively than do FV tumors without malignant US features [40,41].

Recently, encapsulated FV tumors were classified into invasive and noninvasive encapsulated FV PTC [42]. Noninvasive encapsulated

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**Fig. 3.** Diffuse sclerosing variant of papillary thyroid carcinoma in a 22-year-old male patient.

A, B. Transverse (A) and longitudinal (B) ultrasonography shows an ill-defined hyperechoic nonmass-like lesion (arrows) with scattered microcalcifications in the left thyroid gland. Metastatic lymph nodes were noted in the left lateral neck (not shown).
FV tumors have a very low risk of adverse outcomes, whereas invasive encapsulated FV tumors have a 12% risk of recurrence and a 2% risk of tumor-related mortality. It has been proposed that the noninvasive encapsulated FV be termed noninvasive follicular thyroid neoplasm with papillary-like nuclear features (NIFTP) [42]. NIFTP occupies an intermediate position between true benign lesions and invasive cancer and can be removed by limited surgery. Surgical removal is necessary to evaluate the extent of tumor invasion and to prevent possible progression to invasive encapsulated FV PTC [43]. Therefore, the pathological evaluation of these tumors should include the microscopic examination of the entire tumor capsule to exclude invasion.

A multicenter study found that NIFTP lacked malignant US features and frequently showed hyperechogenicity or isoechogenicity, a circumscribed margin, and no calcification on US. It recommended the performance of US-guided core needle biopsy rather than US-guided fine needle aspiration for better triage and to facilitate surgical management. US evaluation is pivotal in determining the next step of FV PTC management. In a previous study, central nodal metastasis was identified in only one of 34 NIFTP cases (2.9%), none of which exhibited distant metastasis or recurrence [44,45]. Infiltrative FV PTCs are not overlooked when the current system for detecting thyroid nodules based on US features is used. However, differentiating between encapsulated FV tumors and NIFTPs via US is still challenging and requires further research (Fig. 4).

Cribriform-Morular Variant
The CMV is a rare type of tumor that occurs in patients with autosomal dominant inherited familial adenomatous polyposis (FAP), although sporadic isolated cases have been reported [46,47]. Multiplicity usually is present in patients with FAP [48]. In such patients, CMV PTC may be diagnosed before colonic manifestations are apparent; hence, its proper identification is essential for the prompt assessment of colonic disease and APC gene status, as well as timely familial counseling. CMV PTC has an early age of onset (age range, 19 to 46 years) and female predominance [48]. It is histologically characterized by the papillary growth of tall columnar cells, a cribriform pattern without colloids, and the presence of spindle cells, squamoid morules, and nuclear clearing. β-Catenin immunostaining is important for the definitive diagnosis of CMV PTC [49].

Previous studies have shown that that most CMV tumors do not exhibit malignant features on US [48]. Rather, their most common US feature is the presence of oval, circumscribed, heterogeneous, and hypoechoic solid nodule(s) without a hypoechoic halo or calcification (Fig. 5). It is not easy to diagnose this type of PTC preoperatively based solely on US features and cytology.

Warthin-like Variant
First described in 1995, the WV is a rare entity that resembles the Warthin tumor of the salivary gland [50]. It consists of cells with oncocytic cytoplasm and the nuclear features of papillary carcinoma that line the papillary structures with dense lymphocytic infiltration in the stalks. The presence of lymphocytic stroma in a thyroid tumor is thought to indicate a favorable prognosis, with tumor-associated lymphocytes contributing to the control of tumor growth [50]. Associated Hashimoto thyroiditis was observed in 93% to 100% of the reported patients with WV PTC [51]. The WV has a significantly

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**Fig. 4.** Noninvasive follicular thyroid neoplasm with papillary-like nuclear features (NIFTP) in a 44-year-old man. A, B. Transverse (A) and longitudinal (B) ultrasonography shows an oval smooth isoechoic nodule (arrows) in the right thyroid gland.
Conclusion

The US features at the time of diagnosis are helpful in predicting the biological behavior of PTC. With active surveillance replacing immediate biopsy or surgery for indolent, small PTCs, an understanding of the US characteristics of the variants of PTC will

lower rate of BRAF mutations than does classic PTC (65% vs. 84%).

WV PTC has a favorable prognosis but can be misdiagnosed as a probably benign nodule or focal thyroiditis on US. All cases are associated with heterogeneous parenchyma in the background [52]. Table 2 is summerized in the incidence and common US features of PTC variants.

| Variant                        | Incidence | Common ultrasonographic feature                      |
|-------------------------------|-----------|------------------------------------------------------|
| Tall cell variant             | 4%–17% of PTCs | Typical malignant features*, frequent nodal metastasis |
| Columnar cell variant         | No report | Typical malignant features vs. circumscribed border   |
| Hobnail variant               | No report | Typical malignant features, frequent nodal metastasis |
| Solid variant                 | 3% of PTCs | Typical malignant features vs. circumscribed border   |
| Diffuse sclerosing variant    | 0.7%–6.6% of PTCs | Ill-defined mass, scattered microcalcifications, nodal metastasis |
| Follicular variant (encapsulated) | 10%–20% of all thyroid cancers | Solid hypoechoic or isoechoic nodule |
| Cribriform-morular variant    | 1% of FAP patients | Solid oval to round nodule without calcification       |
| Warthin-like variant          | No report | Benign-looking nodule, heterogeneous parenchyma        |

PTC, papillary thyroid carcinoma; FAP, familial adenomatous polyposis.

*Typical malignant features include a solid hypoechoic nodule with microcalcification, a spiculated/microlobulated margin, or nonparallel orientation.
Variants of papillary thyroid cancer

facilitate the individualized management of patients with PTC.

ORCID: Jung Hee Shin: http://orcid.org/0000-0001-6435-7357

Conflict of Interest
No potential conflict of interest relevant to this article was reported.

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