Ultrasound Imaging of Thyroid Pathologies: A Pictorial Review

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Abstract

With its complex embryological origin, the thyroid can be affected by various congenital, developmental, benign, and malignant pathologies. Ultrasound, which is free from radiation and offers good spatial resolution, is the initial modality of choice in congenital hypothyroidism and is used in conjunction with scintigraphy. High-resolution ultrasound also aids in early diagnosis, risk stratification, and follow-up of nodules. While fine-needle aspiration cytology is the preferred method for further evaluation of thyroid nodules, ultrasound guidance reduces the likelihood of obtaining nondiagnostic samples. Numerous risk stratification guidelines for thyroid nodules have been developed by various societies over the past decade, with the most popular being the American College of Radiology—Thyroid Imaging Reporting and Data System. A comprehensive understanding of the varying morphological appearances of thyroid nodules and the consistent use of risk stratification guidelines can accurately detect incidental malignancies while avoiding unnecessary intervention in seemingly benign nodules.

Keywords
- thyroid nodules
- TI-RADS
- colloid nodules
- papillary carcinoma
- thyroglossal duct
- thyroid dysgenesis
- thyroiditis

Introduction

Thyroid disease is one of the most common endocrine diseases, with an estimated prevalence of approximately 42 million people suffering from thyroid disease in India.1 The disease processes can be congenital, developmental, inflammatory, or neoplastic, affecting both children and adults. Thyroid nodules are common in women and the elderly and are attributed to hormonal, environmental, genetic, and hereditary factors.2 The incidence of thyroid nodules ranges from 20 to 76%, with increasing diagnosis of incidental nodules on high-resolution ultrasonography.2 Fine-needle aspiration cytology (FNAC) is a cost-effective and minimally invasive test of choice in establishing a definitive diagnosis of these nodules. Malignant disease is, however, seen in only 5 to 10% of thyroid nodules.3 Hence, various imaging based TIRADS (Thyroid Imaging Reporting and Data System) guidelines have been devised for risk stratification of the nodules and reduction of unnecessary FNACs. This pictorial review gives an overview of ultrasound imaging techniques, imaging anatomy, developmental, benign, and malignant thyroid pathologies, and imaging guidelines for reporting of thyroid nodules.

Embryology and Anatomy of Thyroid Gland

The thyroid is a shield-shaped organ in the anterior aspect of the neck (“Thyreos” means shield in Greek).4 During the 4th week of intrauterine life, a small diverticulum develops from
the ventral wall of the primitive pharynx in the midline. This primitive gland grows and migrates to the lower part of the neck coursing anterior to the hyoid and larynx. The tract between the pharynx and the migrated thyroid, called the thyroglossal duct, normally involutes at a later stage and only a small pit called foramen cecum remains in the dorsum of the tongue (►Fig. 1). The primitive gland gives rise to the follicular cells, which secrete thyroid hormones. Parafollicular C cells, which secrete calcitonin, arise from the neural crest cells, which migrate into fourth and fifth pharyngeal pouches forming the ultimobranchial body. Migration completes by the 7th week of intrauterine life, and the production of thyroid hormone starts from the 12th week of gestation itself.

The fully developed thyroid gland comprises two lobes, connected by the isthmus at the level of second and third tracheal rings. It is located in the visceral space of the neck, related to the trachea and prevertebral space posteriorly, and to the sternohyoid and sternothyroid muscles anteriorly.

**Ultrasound Imaging Techniques in the Evaluation of Thyroid**

Being a superficial structure, ultrasound is considered the imaging modality of choice in the evaluation of thyroid pathologies, characterization of nodules, and calculation of thyroid volume. High-resolution (7.5–13 MHz) linear array transducers are generally preferred, while lower frequency transducers can be used for larger nodules and masses. The patient lies in the supine position, with a pillow under the shoulder to facilitate hyperextension of the neck. The thyroid is evaluated for nodules, echogenicity, and vascularity. The normal thyroid gland is well-defined, shows uniform homogeneous echotexture, is more hyperechoic than adjacent muscle, with each lobe measuring approximately 4 to 6 cm craniocaudally (CC) and 1.3 to 1.8 cm in the anteroposterior (AP) and transverse (T) dimensions. The volume of each lobe is calculated using the ellipsoid formula \((\text{AP} \times \text{T} \times \text{CC})^{0.52}\) and ranges from 10 to 20 mL in adults. Color Doppler imaging, at a low flow setting, is used to assess the vascularity of the gland as well as nodules. Elastography is a frequently used adjunct technique which measures the hardness of tissues and, thereby, can help to differentiate benign and malignant nodules. There are two types of elastography techniques used in routine practice for thyroid nodules: (1) strain elastography, which depends on the deformation of tissues in response to mechanical stress (manual compression by the probe or using carotid artery pulsations as an internal compression source or by focused ultrasound waves in acoustic radiation force impulse (ARFI) imaging) and (2) shear wave elastography (SWE), which tracks the attenuation of acoustic pulses called shear waves traveling perpendicular to the direction of the ultrasound beam. Contrast-enhanced ultrasound (CEUS) is also an additional technique that demonstrates the tumor microvascularization by intravenous injection of microbubbles (Sulfur Hexafluoride,
SonoVue, 1.2–4.8 mL followed by saline flush). The degree of enhancement (more, less, or equal to normal thyroid), pattern (centripetal or centrifugal or scattered), homogeneity, wash-in and wash-out times, and other features like ring enhancement help in the differentiation of benign and malignant nodules.9

Thyroid Pathologies
Congenital and Developmental

Thyroid Dysgenesis
Dysgenesis of the thyroid includes “ectopia” due to abnormal migration of the gland, “agenesis or athyreosis” where the gland is absent, and “hypoplasia” when it is not optimally developed. Normal thyroid volume (both lobes) ranges from 0.3 to 4 mL in neonates and infants to 1.7 to 6.4 mL by 6 to 8 years and 3 to 8.7 mL by 9 to 11 years of age and is slightly more in girls.10 Ectopic thyroid tissue can be found anywhere along the course of the thyroglossal duct, most commonly (90%) at the base of the tongue (► Fig. 2). Thyroid scintigraphy is the gold standard for the evaluation of children with congenital hypothyroidism. The thyroid may appear normal on ultrasound in hypoplasia, but scintigraphy can still show poor uptake.11 Occasionally, one or more cysts can also be found in the region of the hypoplastic or absent thyroid (► Fig. 3). They are hypothesized to be remnants of the ultimobranchial body or the thyroglossal duct.12

Lack of Obliteration of Thyroglossal Duct
Patency of the thyroglossal duct can lead to the formation of cyst, sinus, or fistula. Thyroglossal duct cyst is the most common congenital neck mass (► Fig. 2). The thyroglossal duct loops in between the developing hyoid bone and thyroid cartilage. This creates a tail of the cyst coursing beneath the hyoid, a feature that helps to differentiate it from other neck cysts. Thyroglossal ducts cysts are seen in the midline, occasionally paramedian, and can be suprathyroid, at the level of hyoid or commonly infrathyroid.13 Thyroglossal fistulas (► Fig. 4) can open at the foramen cecum above till the lower part of the neck below with numerous side branches. The persistent tract may also form ramifications within the hyoid bone or its periosteum.14

Benign

Simple or Hemorrhagic Cyst
Cysts are thought to occur due to an ischemic episode that creates necrosis. This can happen congenitally, developmentally, or due to tumors, most commonly in papillary thyroid
carcinoma. A simple cyst without any septation or solid component is considered to be benign (►Fig. 5).

Benign Follicular Nodule
It includes colloid nodules, nodular goiter, hyperplastic nodules, nodules in Grave’s disease, and macrofollicular subtype of follicular adenoma. All such nodules have a varying composition of follicular cells and colloid within—with more colloid in colloid nodules and more follicular cells and varying fibrosis in hyperplastic nodules (►Figs. 5 and 6). Calcification of inspissated colloid is responsible for the comet tail artifacts, seen on ultrasound.

Thyroiditis
Diffuse inflammation of the thyroid (thyroiditis) can be classified into acute infectious, autoimmune (Grave’s and Hashimoto), subacute granulomatous (De Quervain’s) thyroiditis, and fibrous (Riedel’s) thyroiditis. The thyroid becomes diffusely heterogeneous and hypoechoic (due to lymphocytes) with increased vascularity and micronodules, particularly in Hashimoto thyroiditis (►Fig. 7). There is also an increased risk of primary thyroid lymphoma and papillary carcinoma in Hashimoto thyroiditis.16 Occasionally, thyroiditis can itself be focal, accounting to approximately 5.3% of all thyroid nodules.17 Differentiation between different types of thyroiditis or nodular thyroiditis from other follicular nodules on ultrasound is not possible due to overlapping and variable imaging appearance.16,18

Follicular Adenoma
It is a benign encapsulated neoplasm made up of follicular cells. Follicular adenomas and carcinomas cannot usually be differentiated on ultrasound or FNAC; a biopsy is needed to assess the vascular invasion in carcinomas. However, recent studies suggest that the presence of nodules within nodule appearance along with calcifications is more likely in follicular carcinomas.19

Malignant
Papillary Carcinoma
It is the most common thyroid malignancy (80%) and includes cells arranged in the form of papillae or monolayers. Psammoma bodies and cystic changes are typically associated with them, which correspond to microcalcifications and anechoic areas on ultrasound in the nodule and their nodal metastases (►Fig. 8).7 The follicular variant appears larger and shows more benign sonographic features.20 It has a very favorable prognosis when treated early, especially in patients who are below 30 years of age.15

Follicular Carcinoma
It constitutes 11% of thyroid malignancies. Here, the neoplastic follicular cells show capsular or vascular invasion and have a propensity for hematogenous spread, most commonly to bone and lungs (►Fig. 9). Hurthle cell carcinoma is a variant of follicular carcinoma with oncocytic cells, which has an aggressive course and poor prognosis.7
**Medullary Carcinoma**
It constitutes 4% of thyroid malignancies with an intermediate prognosis.⁷ Arising from the C cells, this tumor produces calcitonin, and it cannot take up iodine. It is most commonly (80%) sporadic. Twenty percent can be familial—either related to MEN 2a, MEN 2b, or non-MEN. Compared with papillary carcinomas, they are larger, more heterogeneous, and vascular on ultrasound (►Fig. 10).²¹

**Anaplastic Carcinoma**
It is a rare thyroid malignancy with aggressive local invasion, amounting to only up to 2% of the total (►Fig. 10). The malignant cells are undifferentiated; hence cannot take up iodine and are unsuitable for radioablation.⁷ Most of the patients have a history of long-standing goiter, as the tumor arises from the transformation of well-defined carcinoma or adenoma. It shows rapid growth and compressive symptoms.¹⁵

**Lymphoma**
Primary lymphoma is rare (1 to 5% of thyroid malignancies). Hashimoto thyroiditis is a predisposing factor. It can be nodular, diffuse, or mixed (►Fig. 10). The nondiffuse types are markedly hypoechoic and have been referred to as pseudocysts. Lesions can be multifocal, hypervascular, and calcification is uncommon.²²

**Metastasis**
It is rare and occurs in patients who already have widespread metastasis in other organs. The most commonly reported tumors metastasizing to the thyroid include lung, breast, renal malignancies, and melanoma.²³

**Evaluation of Nodules and Risk Stratification**
Since 2009, various risk stratification algorithms (TIRADS) have come up for thyroid nodules, of which the most commonly used one is that of the American College of Radiology (ACR).²⁴²⁵ Point scores are given for each morphological feature (►Fig. 11) and the final score and size are used to decide the need for FNAC or follow-up (►Table 1).

**Evaluation of Thyroid Nodules on Ultrasonography**
Composition: Nodules can be solid, mixed solid cystic, entirely cystic, or spongiform. Among these, solid nodules have the highest risk of malignancy.²⁶ Spongiform nodules, characterized by multiple microcystic areas forming more...
**Fig. 5** Benign thyroid nodules. TR1—Completely cystic nodule with dependent colloid (asterisk in A), TR 1—spongiform nodule with more than 50% cystic spaces (B), comet tail artifact (arrow in C), TR 2—solid cystic isoechoic nodule (D), TR3—solid cystic isoechoic nodule with macrocalcification (arrow in E), and TR 3—solid nodule more echogenic than the adjacent parenchyma (asterisk in F). All the nodules were colloid or nodular goiters by FNAC (Bethesda II).

**Fig. 6** Benign thyroid nodules. TR3—Solid isoechoic nodule with peripheral vascularity (A), soft on elastography (B), TR 4—Solid isoechoic nodule with macrocalcifications (arrow in C), TR 4—Solid isoechoic with interrupted rim calcification (arrow in D), TR 4—Solid nodule hypoechoic compared with the adjacent parenchyma (asterisk in E), and TR 4—Solid hypoechoic nodule with rim calcification (arrow in F). FNAC proved them to be nodular goiters (Bethesda II).
Fig. 7  Thyroiditis. Transverse ultrasound view showing diffusely heterogeneous echotexture of the thyroid with rich internal vascularity (A). Solid hypoechoic nodule (TR 4) in a background of thyroiditis seen as diffuse heterogeneous parenchymal echotexture (asterisk in B). Similar nodule with rich internal vascularity in another patient (C and D). FNAC proved them to be lymphocytic thyroiditis (B) and Hashimoto thyroiditis (C and D).

Fig. 8  Papillary carcinoma. Punctate echogenic foci (arrows) within hypoechoic (A), isoechoic—almost entirely solid (B) and solid cystic—isoechoic (C) nodules (TR5, TR4, and TR 4, respectively). (D) A very hypoechoic nodule as compared with the strap muscle (asterisk), with lobulated margins (TRS) and figure (E) shows a hypoechoic nodule with irregular margins (arrowheads) and punctate echogenic foci (TRS), which is hard on elastography (F). All the nodules were diagnosed as papillary carcinomas (Bethesda VI) on FNAC.
Fig. 9  Follicular neoplasms. A solid hypoechoic nodule (TR 4) with rich vascularity diagnosed to be a follicular neoplasm by FNAC (asterisk in A [Bethesda IV]). Another solid hypoechoic nodule with microcalcifications (arrow in B) and lobulated margins (TR 5) diagnosed as follicular variant of papillary carcinoma on FNAC (Bethesda VI), finally proven to be noninvasive follicular thyroid neoplasm with papillary features (NIFTP) on histopathology post thyroidectomy. A TR 4 nodule in left lobe (C) on ultrasound, corresponding to a hypoenhancing nodule (asterisk in D) on CECT, with lytic expansile blow out metastasis of rib and humerus (arrows in E and F), proven to be a follicular carcinoma on histopathological examination.

Fig. 10  Medullary carcinoma (A and B). A solid taller than wide hypoechoic nodule (asterisk in A) with irregular margins, microcalcifications, and extra thyroid extension (arrow in A) into strap muscles (TR 5), associated round hypoechoic cervical lymph nodes (arrow in B), proven to be medullary carcinoma on FNAC (Bethesda VI). Lymphoma (C). A heterogeneous very hypoechoic nodule (asterisks) with lobulated margins (TR 5) with thyroiditis changes in the background parenchyma (arrow in C), proven to be diffuse large B cell lymphoma on biopsy. Anaplastic carcinoma (D–F). Enlargement of left lobe of thyroid by a large TR 4 nodule (D) with internal cystic change and macrocalcification (arrow in E). Another patient with an ill-defined hypoechoic nodule showing extra thyroid extension into trachea (arrowheads in F).
than 50% of the nodule, have a high negative predictive value for malignancy.\textsuperscript{27}

Echogenicity: Nodules are iso, hypo, or hyperechoic if they are equal to, less than, and more than the echogenicity of the normal thyroid. Nodules are very hypoechoic if they are less echogenic than the strap muscles.\textsuperscript{28}

Margins: They can be smooth, ill-defined (more than 50% contour is not made out), lobulated, irregular, or can show overt extrathyroid extension (\textit{\textsuperscript{►}Fig. 10A}) into strap muscles, trachea, esophagus, or larynx, which has the maximum risk of malignancy. The presence of only capsular bulge, border abutment, or loss of echogenic border is considered as minimal extrathyroid extension, the clinical significance of which is controversial.\textsuperscript{24}

Shape: Taller than wide orientation (AP dimension greater than T) happens due to the nonuniform growth of malignant cells in different directions. When this feature is combined with solid composition, there is 93% specificity for malignancy.\textsuperscript{29}

Calci\textsuperscript{c}fications: Dystrophic macrocalcifications with posterior shadowing are commonly seen with benign multinodular goiters. Punctate echogenic foci, which do not show posterior shadowing, correspond to Psammoma bodies histologically and are more commonly seen in malignant nodules.\textsuperscript{30} Comet tail artifacts (\textit{\textsuperscript{►}Fig. 5C}) are echogenic foci with additional posterior reverberation in a triangular pattern, which are strongly associated with colloid nodules.

Vascularity: Neoplasms, hyperplasia of follicles, and granulation tissue in colloid nodules can show vascularity.
However, few papillary carcinomas with dense fibrosis may show poor internal vascularity. Power Doppler, more useful than color Doppler, is angle independent, reduces the artifacts and can also help in an accurate depiction of vascularity from small vessels.\(^{31}\)

**Elastography**

Strain elastography (SE) by compression: In SE, the stiffness of the tissue is displayed as a color spectrum from red (soft), green (medium), to blue (hard). Qualitative scores (►Fig. 12) have been described by Asteria et al (4 points, score 3, or more favoring malignancy) and Rago and Vitti (5 points, score 4, or more favoring malignancy), in the order of increasing stiffness of the nodules.\(^{8,32,33}\) Semiquantitative evaluation can also be done using strain ratio, comparing the stiffness between the thyroid nodule and surrounding normal parenchyma or muscle. A strain ratio of greater than 1.21 has been found to be the best cut-off in differentiating benign and malignant nodules.\(^{34}\)

SE by Acoustic radiation force: Qualitative ARFI imaging (►Figs. 13D and 13F) displays tissue stiffness in the form of a grayscale, darker the nodule, more the stiffness. They have been classified into six grades (grade 4 or more favoring malignancy) based on the increasing proportion of darker/blacker areas by Xu et al.\(^{8,35}\) It was also observed that malignant nodules, owing to their infiltrative margins, show larger areas on elastography images than conventional ultrasound. Nodules were more frequently malignant when the area ratio (area on elastography to that in conventional ultrasound) was greater than 1.09.\(^{36}\)

Point SWE: In point SWE, the stiffness of tissues is depicted in m/s of the shear wave speed inside a region of interest (ROI) of fixed size (►Figs. 13E and 13G). The median shear wave velocity of normal thyroid parenchyma ranges between 1.2 and 3.6 m/s.\(^{37}\) The cut-off between benign and malignant nodules with the best diagnostic efficacy ranges between 2.48 and 2.55 m/s.\(^{38}\) Diffuse thyroid diseases like Hashimoto or Grave’s disease may also show high shear wave values of 2.68 ± 0.5 m/s.\(^{8}\) However, the range of elasticity values that can be depicted is narrow in m/s (usually 0.5–8.4 m/s or 0–10 m/s, depending on the manufacturer) and hence, very hard or very soft tissues are expressed as nonmeasurable (“X.XX m/s”).

Two-dimensional SWE: This is available in single shot or real time modes (►Figs. 6B, -8F, and 13C) by different vendors. It provides tissue stiffness information in a larger area of user selected box and allows quantitative assessment from multiple ROIs drawn within it. Values are expressed either in m/s (shear wave velocity) or kPa (elasticity), and the color scale is graded from blue (soft) to red (hard).\(^{8}\) The elasticity values for normal thyroid parenchyma ranges between 10 and 40 kPa. An elasticity value of 65 and 66 kPa in the nodule or an elasticity ratio between the nodule and surrounding normal thyroid of more than 3.7 is considered as the cut-off for malignancy.\(^{37}\) The values are higher in papillary carcinomas due to their calcifications and fibrous component compared with follicular carcinoma which has more cellular components.\(^{38}\)

A few pitfalls of elastography include lower stiffness in malignancies with necrosis, cystic areas, and smaller sizes, higher stiffness in benign nodules with calcifications or thyroiditis with more fibrous components, dependence on the application of adequate probe compression, need for including adjacent normal thyroid parenchyma in the field for strain or elasticity ratios which may not be possible in large nodules and varying cut-off values of elastography.\(^{39}\)
parameters by different studies to distinguish benign and malignant nodules.

**Contrast-Enhanced Ultrasonography**

Benign nodules: The patterns of enhancement described in benign nodules (► Figs. 14 and 15) include (1) uniform homogeneous or scattered enhancement; (2) iso or hyper-enhancement compared with surrounding normal parenchyma; (3) fast wash-in and slow wash-out; (4) complete hyper- or hypo-enhancing peripheral ring; and (5) no enhancement in cysts. The first three features are related to the homogeneous architecture of benign nodules with a relative lack of necrosis or calcifications as well as preserved vascularity within them. Follicular adenomas can show hyperenhancement due to their rich vascularity. A hyper or hypo-enhancing rim can be seen due to compression of surrounding thyroid parenchyma and interstitial edema, inflammatory exudates, or the presence of cystic areas within the nodule decompressing the pressure on adjacent parenchyma. Benign nodules with necrosis or a higher proportion of cystic areas, however, may show heterogeneous enhancement.

Malignant nodules: In malignancy, necrosis and tumor embolus formation happens within vessels when the growth of the tumor tissue exceeds the vascular supply, which causes hypoenhancement. They also have a heterogeneous composition with varying areas of fibrosis, necrosis, and calcifications within. The enhancement patterns described in CEUS in malignant nodules include (► Figs. 14 & 15): (1) hypo-enhancement with nodule to peri-nodule peak intensity ratio less than 0.9, (2) heterogeneous enhancement, (3) centripetal enhancement attributed to predominant fibrosis in the central parts, (4) slow wash-in, rapid wash-out ratios and time to peak intensities, although follicular

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**Fig. 13** Elastography of thyroid nodules. Benign nodular goiters showing a qualitative strain elastography score of 2 (A and B), appearing soft on SWE with elasticity of 3.61 kPa (C). Another benign nodular goiter evaluated by qualitative virtual touch imaging (VTi, Siemens), appearing similar to the background parenchyma (asterisk) on gray scale, with mean shear wave velocity of 1.8 m/s on point SWE (D and E). Biopsy-proven medullary thyroid carcinoma, appearing blacker than the background thyroid parenchyma (asterisk) on VTI (F), with an area of involvement larger than that in the gray scale image, and mean shear wave velocity values X.XX suggesting that it is higher than the highest possible value of 8.4 m/s (G).
Fig. 14  CEUS enhancement patterns in thyroid nodules. (A and B) Biopsy-proven papillary carcinoma thyroid seen as a solid hypoechoic nodule with microcalcifications (TR5), with heterogeneous centripetal hypoenhancement pattern on CEUS. An incomplete irregular hyperenhancing ring is seen along part of the nodule (arrowheads in B). (C and D) Solid hypoechoic nodule (TR 4) with intranodular vascularity (C), showing homogeneous iso enhancement and a complete regular hyperenhancing ring (arrowheads in D). It was proven to be a nodular goiter by FNAC.

Fig. 15  Schematic diagram of common CEUS enhancement patterns in thyroid nodules. Malignant nodules (A) show heterogeneous centripetal hypoenhancement, irregular hypo- or hyper ring enhancement, and sometimes no enhancement in smaller nodules. Benign nodules (B) show scattered or equal homogeneous enhancement in all areas, which is equal or more than the background parenchyma, hyper- or hypoenhancing complete regular ring, and no enhancement in cysts.
carcinomas with rich vascularity show faster wash-in and slow wash-out times just like follicular adenomas, (5) ill-defined enhancement borders, (6) irregular or incomplete ring enhancement (usually hypo- or sometimes hyperenhancing due to peritumoral immune response), (7) no enhancement, particularly in nodules less than 1 cm, and (8) enhancement area smaller or larger than grayscale image.\textsuperscript{39–41}

Besides aiding in the differentiation of benign and malignant nodules, they can also help in the follow-up of patients postablation or postradioactive iodine therapy by identifying enhancing viable tissue.\textsuperscript{39,42}

**Associated Cervical Lymph Nodes**
Features of malignant lymph nodal metastasis (\textsuperscript{[Figs. 16 and 17]} from the thyroid include\textsuperscript{43} hyperechogenicity, round

![Image of lymph node evaluation](image)

**Fig. 16** Schematic diagram of lymph node evaluation in suspected thyroid malignancy. Suspicious features (A–D), intermediate suspicion (E) when there is loss of echogenic hilum and hilar vascularity, and benign (F) when there is the presence of echogenic hilum and hilar vascularity.

![Image of lymph node metastasis](image)

**Fig. 17** Lymph node metastasis from papillary carcinoma. Longitudinal views of ultrasound show cystic change (asterisk in A), hyper-echogenicity (arrow in B), rich vascularity with lost fatty hilum (C), and microcalcifications (arrows in D). Contrast-enhanced CT images show cystic changes within a large left level V lymph node (arrow in E). Another patient showing calcification within a metastatic node (arrow in F).
shape, microcalcifications, and cystic change. The presence of cystic areas on ultrasonography is more specific for papillary carcinoma of the thyroid. Echogenic foci can be seen in lymph nodes in both medullary carcinoma (due to amyloid) and papillary carcinoma (due to Psammoma bodies). Metastatic lymph nodes are harder with higher elasticity scores on SWE. On CEUS, malignant lymph nodes show poor vascularization/enhancement areas and perfusion deficits due to necrosis, with centripetal enhancement, while benign nodes show iso and centrifugal enhancement.

**Thyroid Nodule Mimic**

Parathyroid adenomas commonly occur along the inferior poles of the thyroid, separated from it by an echogenic capsule, and appear as hypoechoic solid lesions with feeding polar vessels (< Fig. 18 >). In approximately 1 to 7% cases, they can be completely or partially intrathyroid and can be confused with a thyroid nodule. Four-dimensional computerized tomography can help to differentiate between thyroid nodules and parathyroid lesions, as the latter show arterial hyperenhancement, washout, and polar vessels.

Knowledge of the embryological anatomy can help in differentiating complex congenital thyroid pathologies from other neck pathologies. Meticulous application of TIRADS can help to set apart suspicious nodules which require FNAC and those which can be followed up.

**Author Contributions**

S.L.M. was responsible for design, literature search, manuscript preparation; R.G. was responsible for manuscript editing, manuscript review; S.V.C. was responsible for manuscript review; M.R. was responsible for manuscript preparation; and K.V. was responsible for manuscript preparation.

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**Conflict of Interest**

None declared.

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