Endocrine system. THYROID NEOPLASMS

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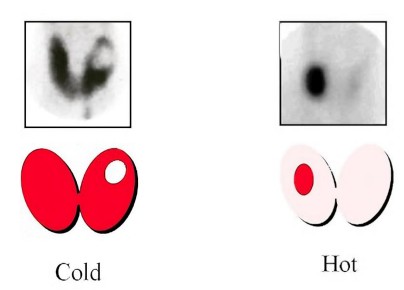
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THYROID NEOPLASMS

- Thyroid tumors range from circumscribed, benign adenomas to highly aggressive, anaplastic carcinomas.
- Fortunately, the overwhelming majority of solitary nodules of the thyroid prove to be either:
- √ benign adenomas.
- ✓ localized, non-neoplastic conditions, e.g.
- dominant nodule in multinodular goiter.
- *simple cysts.
- * foci of thyroiditis.

Benign vs malignant

- Thyroid nodule most likely to be malignant if:
- √ Nodules in younger patients.
- ✓ Nodules in males .
- ✓ Nodules that doesn't take up radioactive iodine in imaging studies (cold nodules).

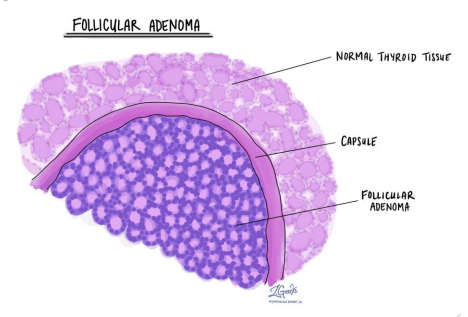


Neoplastic thyroid lesions

- Benign:
- > follicular adenoma.
- Malignant:
- ➤ Papillary carcinoma (accounting for more than 85% of cases)
- ➤ Follicular carcinoma (5% to 15% of cases)
- Anaplastic (undifferentiated) carcinoma (<5% of cases)</p>
- Medullary carcinoma (5% of cases)

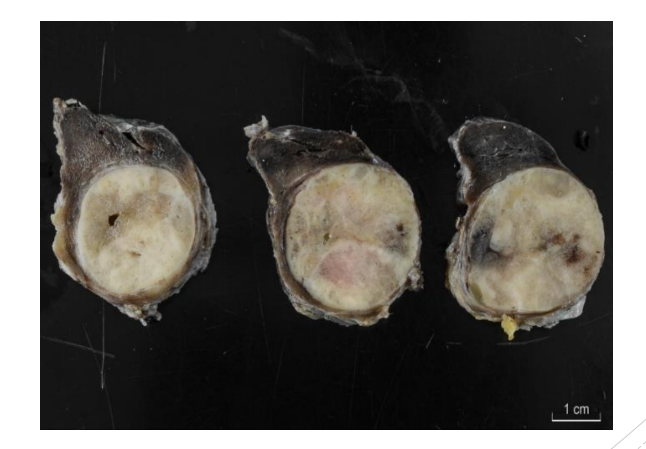
Follicular adenoma

- Adenomas of the thyroid are benign neoplasms derived from follicular epithelium.
- Follicular adenomas usually are solitary, DDX??
- the vast majority of adenomas are nonfunctional, a small proportion produce thyroid hormones (toxic adenomas), causing clinically apparent hyperthyroidism.



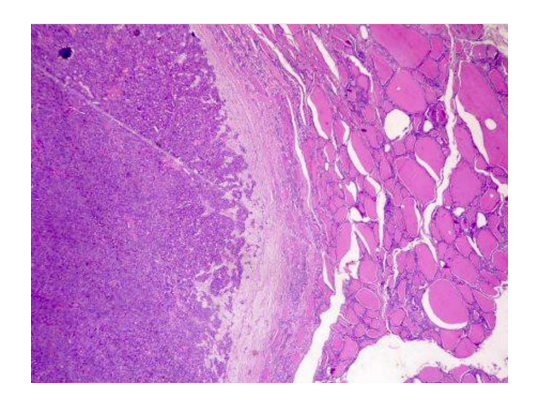
Morphology

• Solitary, encapsulated, variable size (1 - 10 cm).





- Closely packed follicles.
- Completely enveloped by thin fibrous capsule
- surrounding thyroid tissue shows signs of compression.





Lobectomy (not enucleation).

- Carry an excellent prognosis
- > do not recur or metastasize.

Thyroid carcinoma

- 1% of all cancer in U.S., 0.2% of all cancer deaths.
- Increasing incidence due to new diagnostic practices which detect smaller tumors.
- 20 year survival is 90%, because most are indolent papillary carcinomas
- A female predominance has been noted among patients who develop thyroid carcinoma in the early and middle adult years (Often estrogen receptor positive).

1. Papillary Carcinoma.

- The most common types of thyroid carcinoma.
- Female predominance; F:M ratio = ~3:1
- Median age of diagnosis in 50s
- <u>Ionizing radiation</u> is the best established risk factor.
- Mainly 2 genes are involved:
- 1. BRAF amplification.
- 2. RET gene rearrangment .

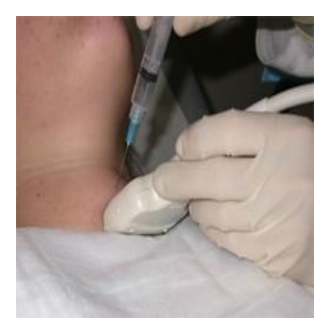
Clinical features.

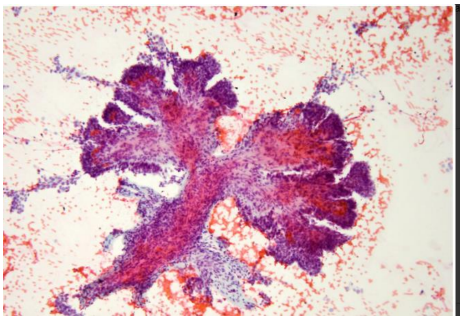
- Presented as Painless palpable thyroid mass.
- The diagnosis is first rendered on ultrasound guided preoperative fine needle aspiration cytology
- Surgical pathology report of a resected specimen provides further information about the subtyping (i.e., variant) and microstaging

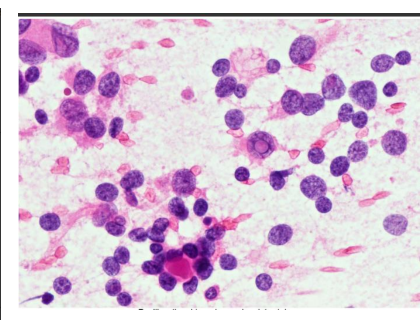
Commonly treated with surgical resection.



ultrasound guided pre-operative fine needle aspiration cytology.

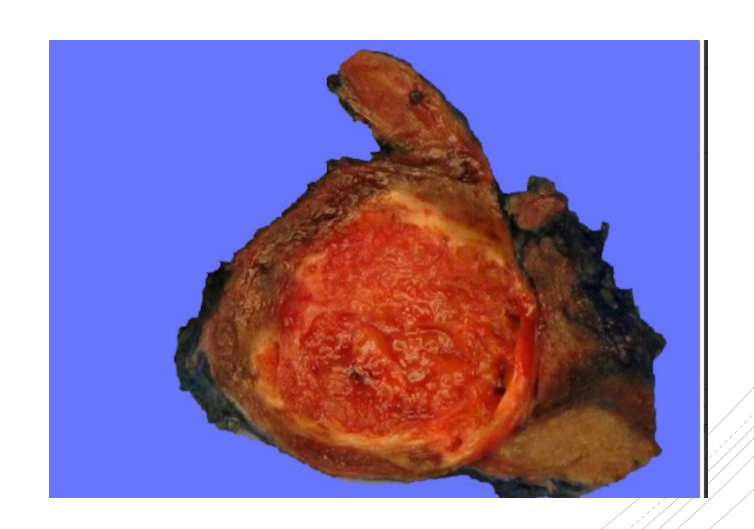


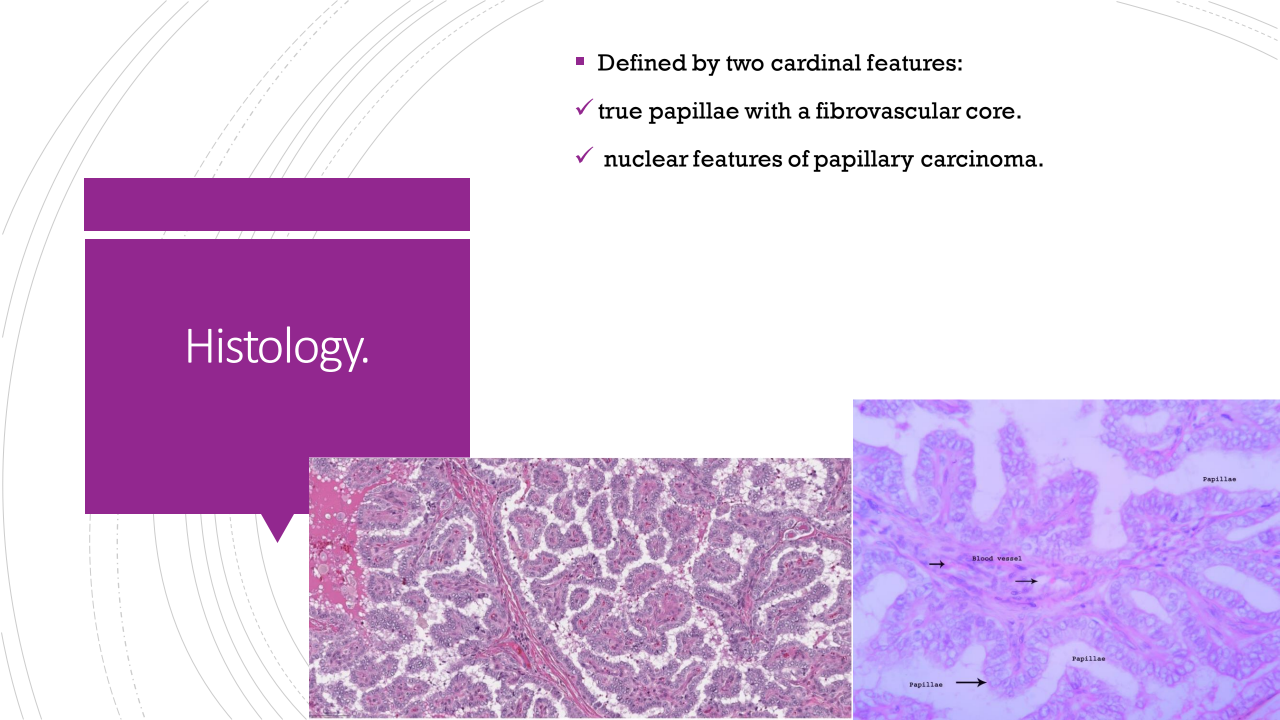


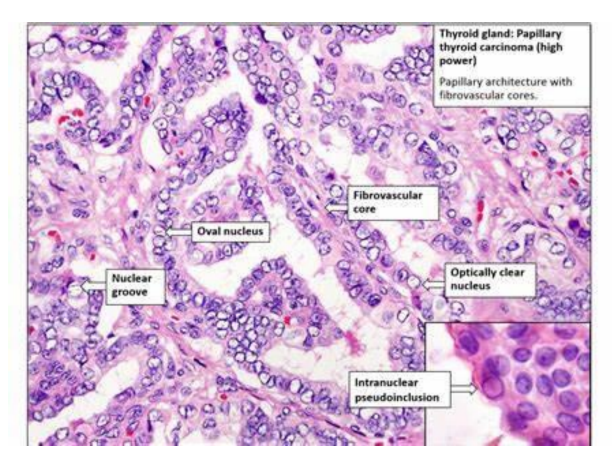


Solid or cystic mass with papillary projections

Morphology

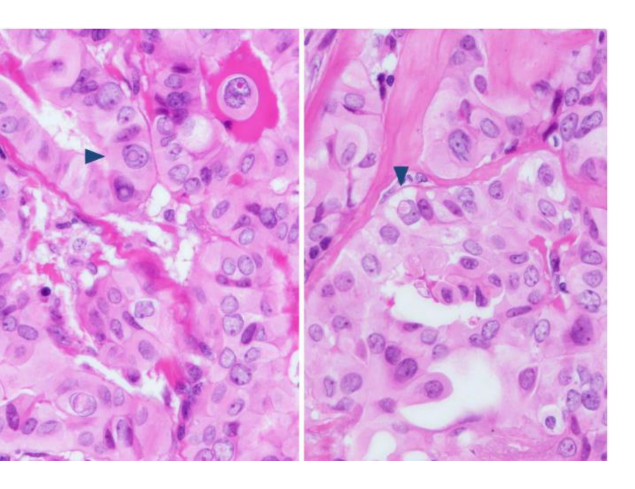


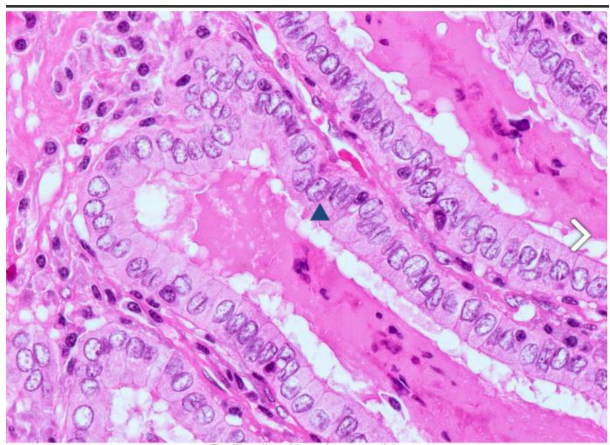






- > irregular nuclear contour.
- > nuclear groove.
- > nuclear pseudoinclusion





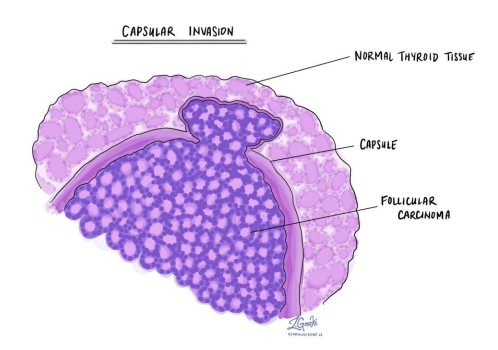
Papillary thyroid carcinoma nuclei:

2. Follicular Carcinoma.

- Thyroid carcinoma with follicular differentiation but no papillary nuclear feature.
- Follicular lesion with capsular or vascular invasion but without papillary nuclear features.
- More common in women and in areas with <u>dietary</u> iodine <u>deficiency</u>.
- The peak incidence between the ages of 40 and 60 years.
- GENETIC FACTORS:
- ✓ Gain-of-function point mutations of RAS and PIK3CA.
- ✓ Loss-of-function mutations of PTEN.

Two types

- 1. Minimally invasive follicular carcinoma
 - With capsular invasion.
 - With vascular invasion
- 2. Widely invasive.



Clinical features

- Usually "cold" on radionuclide scan
- Does not metastasize through lymphatics but does spread to lungs, liver, bone, brain via blood vessels
- Less than 5% with ipsilateral lymphadenopathy.
- Treatment:
- √ thyroidectomy and radioactive iodine
- √ No nodal dissection is needed

Morphology

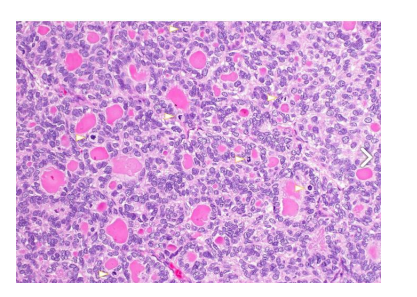
- Tan to brown solid cut surface, can have cystic changes and hemorrhage
- Minimally invasive: usually single encapsulated nodule, with thickened and irregular capsule
- Widely invasive: extensive permeation of capsule or no capsule.

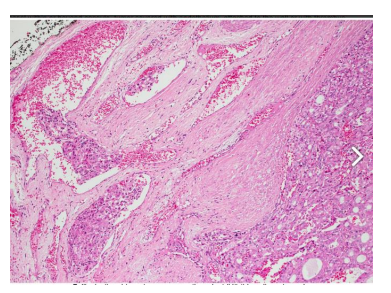


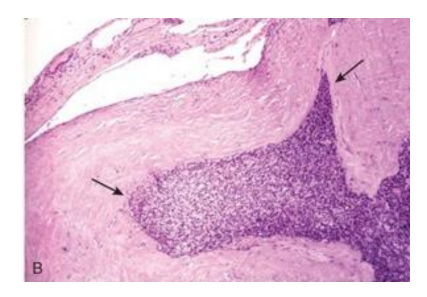




- solid pattern of follicles (small, normal sized or large).
- No nuclear features of papillary thyroid carcinoma
- Invasion of adjacent thyroid parenchyma, capsule (complete penetration) or blood vessels (in or beyond the capsule)







3. . Anaplastic Carcinoma.

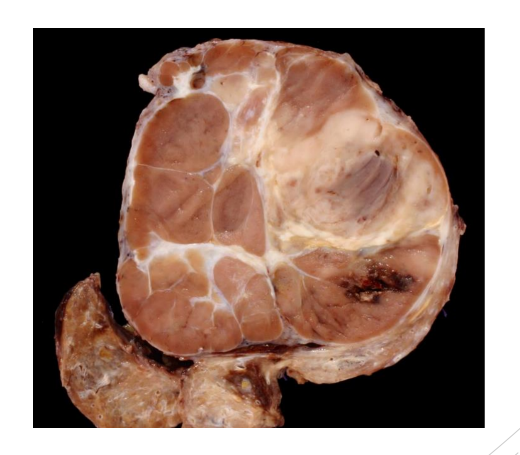
- A highly aggressive thyroid malignancy composed of undifferentiated follicular thyroid cells, devoid of morphologic features of thyroid origin.
- Medium age 60 70 years with incidence to rise with age, F:M = 2:1.
- Higher incidence in areas of <u>dietary iodine deficiency</u>.
- GENETIC FACTORS:
- ✓ Inactivation of TP53.

Clinical features

- Rapidly enlarging, bulky neck mass invades adjacent structures causing hoarseness, dysphagia, dyspnea.
- fixed to the underlying structures.
- Extrathyroidal extension in majority of cases
- Regional nodal metastases and vocal cord paralysis present in up to 40% and 30%, respectively
- Treatment
- Radiation therapy, surgery when feasible or chemoradiation either concurrently or sequentially

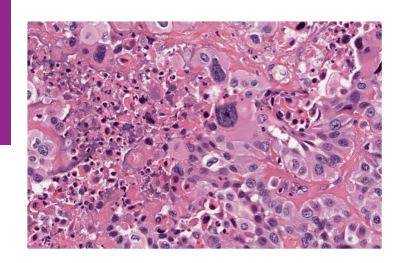
Bulky solid mass (mean: 6 cm) with zones of necrosis or variegated appearance.

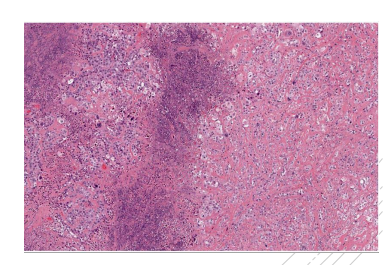
Morphology



Histology

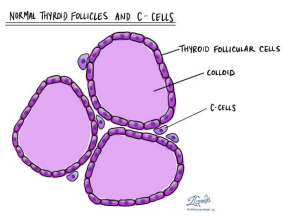
- Common features include:
- ✓ widely invasive growth.
- ✓ extensive tumor necrosis.
- \checkmark marked nuclear pleomorphism .
- ✓ high mitotic activity

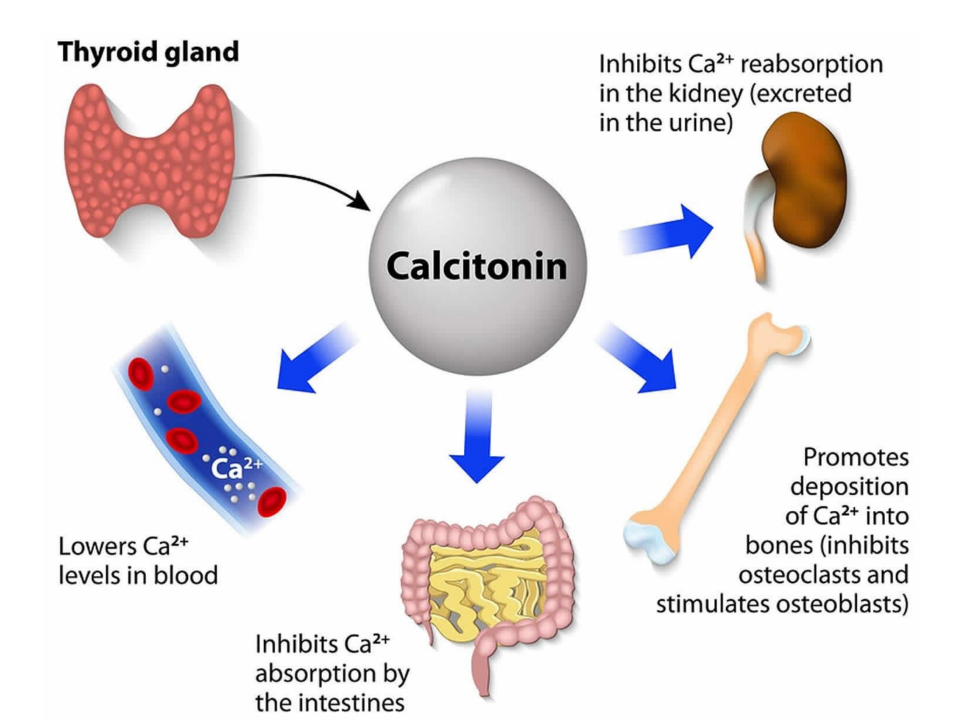


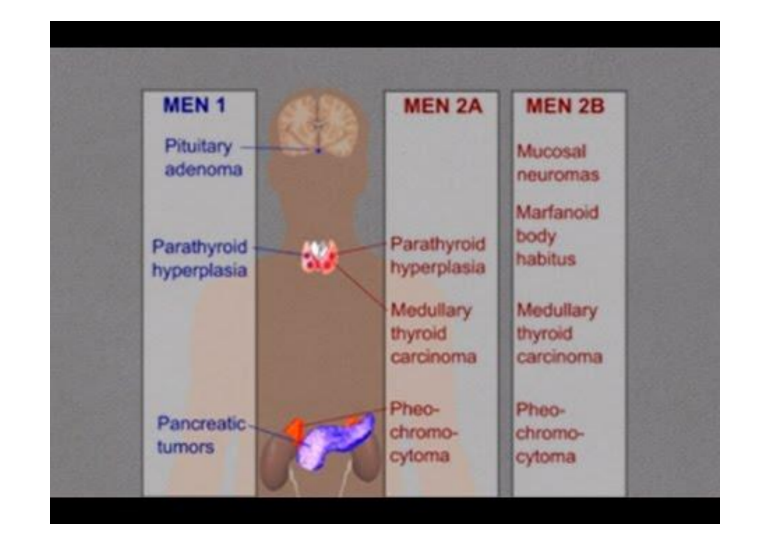


4. Medullary Carcinoma.

- Neuroendocrine tumor derived from C cells (formerly called parafollicular cells), which secrete calcitonin
- 1 2% of thyroid carcinomas
- Either sporadic (nonhereditary) or familial (hereditary)
 - **Sporadic**: 70%, age 40 60, solitary
 - Familial: 30%, younger patients (mean age 35).
 - ✓ Occurring in the setting of MEN syndrome 2A or 2B,
 - ✓ familial medullary thyroid carcinoma without an associated MEN syndrome





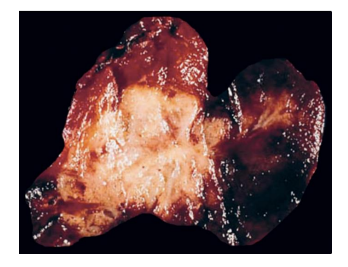


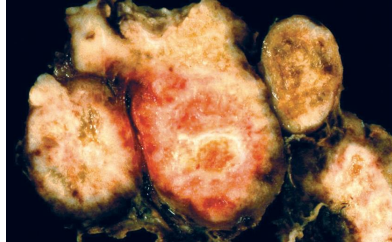
Clinical features

- Presents with painless thyroid mass, cold on scanning
- Up to 75% of patients have nodal metastasis.
- Serum calcitonin correlates with tumor burden.
- Patients with metastasis may have severe diarrhea and flushing
- Some tumors may produce ACTH or CRH (Cushing syndrome).

- **Sporadic**: typically presents as a single circumscribed but nonencapsulated, gray-tan mass
- Familial: generally bilateral / multiple foci.

Morphology







Wide variety of morphology:

Round.
Plasmacytoid.
polygonal
spindle cells.

•Eosinophilic to amphophilic granular cytoplasm due to secretory granules

• Stroma has amyloid deposits from calcitonin

