Mediastinum Tumors

**General**
Region located between the lungs, sternum, spine, thoracic inlet, and diaphragm.

About half of tumors are asymptomatic→ identified on imaging.
Symptoms often result from compression/invasion of structures→ cough, pain, dyspnea.
May block superior vena cava→ “SVC syndrome”→ face swelling, distended neck veins, distended collaterals→ often malignant→ Adults: think lung cancer or lymphoma; Kids: Leukemia/lymphoma.

**Differential Diagnosis by Location:**

<table>
<thead>
<tr>
<th>Anterior</th>
<th>Superior</th>
<th>Middle</th>
<th>Posterior</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thymic tumors</td>
<td>Thymic tumors</td>
<td>Pericardial cyst</td>
<td>Neurogenic tumors</td>
</tr>
<tr>
<td>Germ cell tumors</td>
<td>Thyroid tumors</td>
<td>Bronchial cyst</td>
<td>Schwannoma</td>
</tr>
<tr>
<td>Parathyroid tumors</td>
<td>Lymphoma</td>
<td>Lymphoma</td>
<td>Neurofibroma</td>
</tr>
<tr>
<td>Lymphoma</td>
<td>Parathyroid tumors</td>
<td></td>
<td>Ganglioneuroma</td>
</tr>
<tr>
<td>Paragangioma</td>
<td></td>
<td></td>
<td>MPNST</td>
</tr>
<tr>
<td>Hemangioma</td>
<td></td>
<td></td>
<td>Paraganglioma</td>
</tr>
<tr>
<td>Lipoma</td>
<td></td>
<td></td>
<td>Neuroblastoma</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Gastrointestinal cyst</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Bronchogenic cyst</td>
</tr>
</tbody>
</table>

**The Classic 5 “T’s” of Anterior Mediastinal Masses**
- Thymus
- Thyroid
- Teratoma
- Terrible lymphoma
- Thoracic Aorta

**Developmental Cysts**
Congenital anomalies that develop during embryogenesis.

**Bronchogenic Cyst**
Abnormal tracheobronchial tree branching.

Often well-formed structures resembling bronchus.
Contain a combination of: Ciliated epithelium (1), cartilage (2), submucosal glands (3), smooth muscle, and/or degenerative changes. Unilocular.

Cured by excision. Can get infected.

Can be hard to distinguish from esophageal duplication cysts if ciliated and no cartilage, can say simply “Foregut cyst”

**Gastrointestinal Duplication Cysts**
Attached to the GI tract (but lumens not contiguous, unlike a diverticulum), with epithelium that resembles some part of the GI tract, and a well-developed double layer of smooth muscle (resembling normal bowel layers). NO Cartilage.

**Esophageal Duplication Cyst:** Columnar (ciliated or non-ciliated), squamous, or mixed epithelium. Can contain heterotopic lung or thyroid.

**Enteric Duplication Cyst:** Variable epithelium, usually gastric or duodenal.
Thymic Tumors

**Thymoma**

Thymic epithelial neoplasms with a *variety* of histologic patterns. Overall rare, but most common mediastinal tumor in adults.

**Multiple subclassifications** (see below), but **stage is much more important prognostically**! *(All subtypes can behave aggressively or indolently, mostly important to aid in recognition and DDX)*

Frequent association with **paraneoplastic syndromes**: Most common = **Myasthenia gravis** *(autoantibodies block acetylcholine receptors between muscle & nerves → weakness)*

Other syndromes: Collagen and autoimmune disorders (e.g., lupus), immunodeficiencies, endocrine disorders, dermatologic disorders, enterocolitis, etc..

<table>
<thead>
<tr>
<th>Type</th>
<th>Composition</th>
<th>Proportion Epithelium</th>
<th>Proportion Lymphocytes</th>
<th>Prognosis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Type A</strong></td>
<td>Bland spindled to ovoid cells, few or no admixed lymphocytes</td>
<td>Predominant, spindled/oval</td>
<td>Few/none</td>
<td>Excellent</td>
</tr>
<tr>
<td><strong>Type AB</strong></td>
<td>Both lymphocyte poor (type A) and lymphocyte-rich (type B) components, with a significant proportion of immature T cells</td>
<td>Significant</td>
<td>Significant</td>
<td>Very good</td>
</tr>
<tr>
<td><strong>Type B1</strong></td>
<td>Predominantly lymphocytes with dispersed epithelial cells (that do not form clusters)</td>
<td>Low, no clusters, polygonal</td>
<td>Predominant</td>
<td>Very good</td>
</tr>
<tr>
<td><strong>Type B2</strong></td>
<td>Predominantly lymphocytes, with small clusters of epithelial cells</td>
<td>Low, small clusters</td>
<td>Significant</td>
<td>Fair</td>
</tr>
<tr>
<td><strong>Type B3</strong></td>
<td>Predominantly atypical polygon epithelial cells in sheets.</td>
<td>Predominant, epithelioid</td>
<td>Few</td>
<td>Fair, often high stage</td>
</tr>
<tr>
<td><strong>Micronodular with lymphoid stroma</strong></td>
<td>Multiple small tumors with bland spindled cells surrounded by lymphoid stroma</td>
<td>Significant, spindled</td>
<td>Significant. B &amp; T cells, without epithelial cells</td>
<td>Excellent</td>
</tr>
<tr>
<td><strong>Metaplastic</strong></td>
<td>Biphasic tumor consists of solid polygon epithelial cells in a background of bland spindled cells</td>
<td>Predominant, epithelioid and spindled</td>
<td>Few/none</td>
<td>Very good</td>
</tr>
</tbody>
</table>

**Subtyping:**

Some thymomas are heterogeneous and show multiple patterns of growth. In these cases, list the different patterns quantified by %. Also, be careful definitely subtyping a thymoma on a limited sampling (likely best to just Dx as “Thymoma” and give the pattern(s) present in the biopsy).

Note: Type AB thymomas are inherently heterogeneous.

**Immunohistochemistry:**

Most do not require IHC for subtyping. Often used to differentiate from Non-thymomas.

Thymic epithelial cells $\rightarrow$ AE1/AE3, p63, PAX8.

T-Cells in thymus $\rightarrow$ CD5, CD3, TdT (immature thymic T-cells)
Thymoma (continued)

Adapted from: Twitter @lauraebrown, Laura Brown, MD, UCSF Hematopathology, 2019.

Thymic Tumor

- Lymphoid Component
  - Yes
  - Absent/sparse

- Epithelial Cell Type
  - Epithelioid
  - Spindled

Type A Thymoma

Spindled/oval cells with few or no admixed immature lymphocytes. Bland nuclei with powdery chromatin. Can have a microcystic appearance.

Usually low stage. Often lobulated and circumscribed/encapsulated.

Excellent prognosis.

Type AB Thymoma

2 components: A) lymphocyte-poor spindle cell component and B) lymphocyte-rich component

Varying proportions, but > 10% of tumor with moderate infiltrate of immature TdT+ T-cells.

Usually low stage, lobulated, and very good prognosis.

Type B1 Thymoma

Closely resembles normal thymus: Dispersed epithelial cells that do not form clusters and are set in a dense background of immature T cells mimicking thymic cortex. Also has areas of medullary differentiation (nodular pale areas ± Hassall’s corpuscles; mostly TdT-T cells with a substantial B-cell population).

Usually nodular with a very good prognosis.

Type B2 Thymoma

Epithelioid

- Lymphocytes

Type B3 Thymoma

Thymic Carcinoma

Adapted from: Twitter @lauraebrown, Laura Brown, MD, UCSF Hematopathology, 2019.
Type B2 Thymoma

Polygonal neoplastic epithelial cells set in a background of numerous immature T cells.

Epithelial cells denser than in B1 and are usually clustered with round vesicular nuclei.

Often encapsulated with a fair to good prognosis.

Type B3 Thymoma

Mild or moderately atypical polygonal pink epithelial cells with lobules of sheet-like or solid growth with fibrous septae. Often few intermingled immature T-cells.

Usually poorly circumscribed → extensions into mediastinal fat/organs → most patients have local symptoms (e.g., chest pain or SVC syndrome) → fair prognosis overall, frequent recurrences.

Micronodular Thymoma with Lymphoid Stroma

Multiple epithelial nodules surrounded by prominent lymphoid stroma containing mature B and T cells and devoid of epithelial cells.

May contain germinal centers and/or plasma cells.

Excellent prognosis

Metaplastic Thymoma

Biphasic: Composed of alternating areas of solid epithelial cells and bland slender spindle cells. Absent to few lymphocytes.

Very rare. No paraneoplastic syndrome.

YAP1-MAML2 gene fusions

Microscopic Thymoma: Multifocal thymic epithelial proliferations, < 1mm, composed of bland spindled to polygonal cells in well-circumscribed nodules embedded in the medulla or cortex. Very rare.

Sclerosing Thymoma: Abundant collagen-rich stroma in an otherwise conventional thymoma. Very rare.

Lipofibroadenoma: Benign thymic tumor that resembles a fibroadenoma of the breast. Very rare.
Thymic Carcinomas

Thymic epithelial tumor with **malignant cytologic features** that lacks thymic organization. **Resembles conventional carcinomas in other organs.**

Often unequivocal cytologic atypia. Often **uncapsulated and no** fibrous septae. Variable T cell infiltrate

IHC: (+) CK AE1/AE3, p63, PAX8, CD5, CD117, GLUT1, MUC1. Focal Synaptophysin often.

**Types of Thymic Carcinoma**

**Squamous Cell Carcinoma:** Most common type of thymic carcinoma. Resembles SCC elsewhere. Lacks normal thymic architecture (e.g., lobulation, lymphocytes, etc...). Frankly invasive into nearby structures and often present with symptoms. Often eosinophilic cytoplasm and abundant stroma.

**Basaloid Carcinoma:** High N:C basaloid appearance with cystic and papillary architecture and peripheral palisading. Lots of mitoses and necrosis. Very aggressive.

**Mucoepidermoid Carcinoma:** Like in other organs (Squamoid cells, mucus-producing cells, and intermediate cells). MAML2 translocations.

**Lymphoepithelioma-like Carcinoma:** poorly-differentiated squamous cell carcinoma with an associated rich lymphoplasmacytic infiltrate (resembles nasopharyngeal carcinoma). Often EBV-positive.

**Clear Cell Carcinoma:** Composed predominantly of cells with vacuolated clear cytoplasm.

**Sarcomatoid Carcinoma:** consists completely or partly of spindled cells. If heterologous elements → Carcinosarcoma.

**NUT Carcinoma:** Like elsewhere, NUT gene rearrangement. Monomorphic round cells with characteristic abrupt keratinization. Often stain with squamous markers. NUT IHC +. Extremely aggressive.

**Adenocarcinomas:** Heterogeneous group showing glandular and/or mucin production.

**Undifferentiated Carcinoma**

**Thymic Neuroendocrine Tumors**

Rare. Classified using same criteria as in lung. No smoking association. Can see with MEN1. (See Lung Tumor Notes for more info)

<table>
<thead>
<tr>
<th></th>
<th>Typical carcinoid</th>
<th>Atypical carcinoid</th>
<th>Large cell neuroendocrine carcinoma</th>
<th>Small cell carcinoma</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Mitoses/2mm²</strong></td>
<td>0-1</td>
<td>2-10</td>
<td>&gt;10 (median 70!)</td>
<td>&gt;10 (median 80!)</td>
</tr>
<tr>
<td><strong>Necrosis</strong></td>
<td>No</td>
<td>Focal, if any</td>
<td>Yes</td>
<td>Yes, extensive</td>
</tr>
<tr>
<td><strong>Morphology</strong></td>
<td>Organoid or trabecular growth, uniform polygonal cells, finely granular “salt and pepper” chromatin</td>
<td>Large cell size, vesicular to coarse chromatin, frequent prominent nucleoli, and abundant cytoplasm</td>
<td>Small fusiform to round cells, scant cytoplasm, finely granular chromatin, Lots of mitoses</td>
<td></td>
</tr>
<tr>
<td><strong>Ki-67</strong></td>
<td>Up to 5%</td>
<td>Up to 20%</td>
<td>40-80%</td>
<td>Almost 100%</td>
</tr>
<tr>
<td>Combined with non-small cell component</td>
<td>No</td>
<td>No</td>
<td>Sometimes</td>
<td>Sometimes</td>
</tr>
</tbody>
</table>

**Adapted from:** WHO Classification of tumors of the lung, pleura, thymus, and heart. 2015.

**IHC Markers of Neuroendocrine Differentiation:** Synaptophysin, Chromogranin, INSM1. Less so CD56. Cytokeratins often show perinuclear “dot-like” staining.
**Germ Cell Tumors**

*Note: For more info, refer to the Testicle and Ovary guides*

Morphologically identical to gonadal counterparts!
Associated with Klinefelter syndrome (XXY)
- Prepubertal → Mostly teratomas or Yolk Sac
- Women → Mostly teratomas
- Men → Teratomas, Seminoma, YST, and mixed

**Seminoma**
Large polygonal cells with clear to eosinophilic cytoplasm, distinct cell membranes, vesicular chromatin, and prominent nucleoli. Fibrous septae and nested architecture
Lymphocytic infiltrate; Sometimes granulomas

**Yolk Sac Tumor**
Many patterns/architecture. Often hypocellular myxoid areas
- Most common = reticular/microcystic
- Can also be solid, papillary, etc...
Classic: Schiller-Duval Bodies
Hyaline globules. Elevated Serum AFP

**Embryonal Carcinoma**
Large “Primitive” cells
Vesicular nuclei with prominent nucleoli
Coarse, basophilic chromatin. Amphophilic cytoplasm
Variable architecture (nests, sheets, glands)

**Choriocarcinoma**
Malignant cytotrophoblasts (mononuclear) and syncytiotrophoblasts (multinucleated)
Abundant Hemorrhage

**Teratoma**
Composed of tissues from 2-3 germ layers.
Common elements: Skin (with adnexal structures), Cartilage, GI, Brain, etc... Very good to excellent prognosis.
* Mature → exclusively mature (adult-type) tissues
* Immature → has immature fetal/embryonic tissue

**Germ Cell Tumor Immunohistochemistry:**

<table>
<thead>
<tr>
<th>IHC Stain</th>
<th>Seminoma</th>
<th>Embryonal Carcinoma</th>
<th>Yolk Sac Tumor</th>
<th>Choriocarcinoma</th>
</tr>
</thead>
<tbody>
<tr>
<td>SALL4</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>OCT 3/4</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>D2-40</td>
<td>+</td>
<td>+/-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CD117</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CD30</td>
<td>-</td>
<td>+</td>
<td>-/+</td>
<td>-</td>
</tr>
<tr>
<td>Glypican 3</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>+/-</td>
</tr>
</tbody>
</table>
**Soft Tissue Tumors**

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**Thymolipoma**

Encapsulated tumor with mature adipose tissue and interspersed normal thymic tissue.

Benign → cured with excision. Rare.

**Lipoma**: mature adipose tissue only (like elsewhere). Rare in mediastinum.

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**Liposarcoma**

Similar to liposarcomas in soft tissue.

Most common sarcomas of mediastinum, often well-differentiated liposarcomas or dedifferentiated liposarcomas → both have giant marker and ring chromosomes that contain amplified regions of 12q including MDM2 and CDK4 (detect with MDM2 FISH)

**Well-differentiated liposarcoma**: Range of appearances. Variable lipoblasts and hyperchromatic atypical cells in a background of adipocytes and fibrous tissue.

**Dedifferentiated liposarcoma**: Contain an WDL component, with an abrupt transition to another component, which is usually an undifferentiated pleomorphic sarcoma

Often poor prognosis.

---

**Synovial Sarcoma**

Malignant spindle cell neoplasm of uncertain histogenesis. Poor prognosis.

Like in soft tissue, monophasic or biphasic proliferation of spindled cells with stubby nuclei and frequent Stag-horn vessels.

IHC: Patchy EMA and CK (particularly strong in epithelial areas). Usu. CD99 (+). TLE-1 (+)

Molecular: SS18-SSX gene fusions t(X;18)

---

**Solitary Fibrous Tumor**

Usually benign.

“**Patternless pattern**” of varying cellularity of bland spindled cells with varying amounts of collagenized stroma.

Prominent “Staghorn vessels” (dilated, thin-walled, branching vessels).

Can be hyalinized or myxoid.

IHC: **STAT6 (+)**. Also, CD34, CD99 (+, but variable).

Molecular: **NAB2/STAT6 gene fusion**
SMARCA4-deficient Thoracic Sarcomas

Diffuse sheets of mildly discohesive, relatively monotonous, and undifferentiated epithelioid cells with prominent nucleoli.
IHC: (+) CD34, SALL4, (+/-) CK
Molecular: SMARCA4 mutations (part of SWI/SNF chromatin remodeling complex, like INI-1)

Schwannoma

Composed entirely of well-differentiated Schwann cells. Very low risk of transformation.
Usually solitary and sporadic in posterior mediastinum.
Typically encapsulated.
Alternating compact spindle cells (Antoni A) and hypocellular less orderly areas (Antoni B)
Rows of nuclear palisading → Verocay bodies.
Axons not present in lesion → pushed to periphery.
Hyalinized blood vessels and lymphoid aggregates common.
IHC: Strong, diffuse S100, scattered CD34, moderate calretinin. Neurofilament highlights displaced axons at periphery.

Malignant Peripheral Nerve Sheath Tumor (MPNST)

Malignant. Adults. Frequently in setting of NF1.
Often poor prognosis.
Must arise from a peripheral nerve or pre-existing peripheral nerve sheath tumor or display histologic/IHC evidence of nerve sheath differentiation.
Spindled cells arranged in sweeping fascicles.
Densely cellular areas alternate with less cellular areas giving a “marble-like” effect.
Can have herringbone architecture.
Wavy, buckled nuclei.
Geographic necrosis and/or mitotic activity (often greater than 10/10 HPFs).
IHC: Patchy S100 and SOX10.
Loss of H3K27me3 expression (associated with worse prognosis. Not entirely specific—see with SUZ12 and EED gene inactivation)
**Angiosarcoma**

- **Malignant.** Very aggressive. Typically elderly.
- Variable degrees of vascular differentiation.
- Some areas show well-formed anastomosing vessels, while other areas may show solid sheets of high-grade cells. Can be epithelioid or spindled. Often extensive hemorrhage.
- Unlike benign lesions: significant cytologic atypia, necrosis, endothelial cells piling up, and mitotic figures (although mitoses can be seen in some benign tumors)
- **IHC:** CD31, ERG, FLI1, often CD34

**Sclerosing (fibrosing) Mediastinitis**

- **Non-neoplastic fibrosis** of mediastinum compressing and infiltrating normal structures.
- **Bland spindled cells with lymphoplasmacytic infiltrate**
  - Sometimes dense (keloid-like) collagen.
  - May see dystrophic calcifications.

May be caused by:
- prior infection/response to **Histoplasma** or TB
- **IgG4-related disease**
- Autoimmune diseases
- Radiation

**Neuroblastoma**

- **Most primitive/aggressive**
- **Malignant.** Vast majority <5 years
- SRBCT +/- rosettes, neurofibrillary matrix
- Peripheral neuroblastic tumors derive from the sympathetic nervous system (therefore develop anywhere along the distribution of the sympathoadrenal neuroendocrine system), often in posterior mediastinum.
- Stains: Schwann cells (+) S100, Ganglion cells (+) Synaptophysin, neurofilament

**Ganglioneuroma:** Although some likely represent matured neuroblastoma, it is thought that most are *de novo*. Multiple/diffuse and/or syndrome-related (MEN 2b, Cowden, and NF1) → Ganglioneuromatosis
Thyroid & Parathyroid Tumors

Thyroid Tumors:
Often arise in an extension of the thyroid from the neck (as opposed to ectopic thyroid). Identical appearance, IHC, and behavior to thyroid tumors in the neck (see separate guide).

General IHC:
Tumors derived from follicular epithelium (PTC, follicular carcinoma): (+) TTF1, PAX8, Thyroglobulin, CK
Medullary thyroid carcinoma: (+) TTF1, Synaptophysin, Calcitonin, CK, (-) Thyroglobulin, (+/-) PAX8

Parathyroid Tumors:
Ectopic. Up to 20% of all parathyroid neoplasms are located in the mediastinum (often near/in thymus as they share a common origin in 3rd branchial pouch). Often present with hyperparathyroidism and resulting hypercalcemia (kidney stones, bone pain, etc.). Identical appearance, IHC, and behavior (see separate guide). IHC: (+) CK, Synaptophysin, Chromogranin, GATA-3, PTH. (-) TTF1, Thyroglobulin, Calcitonin; (+/-) PAX8

Lymphomas

Classical Hodgkin Lymphoma:
Most common type of primary mediastinal lymphoma! Peak incidence in late adolescence/young adult. Reed-Sternberg cells (classically large binucleated cells with abundant cytoplasm and prominent nucleoli with perinucleolar clearing) in a background of inflammatory cells. Lacunar RS cells are smaller with hyperlobated nuclei. Often lots of eosinophils.

RS cell IHC: (+) CD30, CD15, MUM1. Characteristic weak PAX5. (-)CD20, CD45

Most common variant: Nodular Sclerosis Classical Hodgkin Lymphoma—cellular nodules separated by dense fibrous bands. Often has lacunar RS cells.

Primary Mediastinal Large B-cell Lymphoma:
Aggressive large B-cell lymphoma arising in the mediastinum. Most often in young adults. Presents with localized mass in thymic area and minimal associated distant lymphadenopathy. Diffuse growth of large cells with abundant, often clear, cytoplasm.

IHC: (+) CD19, CD20, CD79a, PAX5.

Requires clinical exclusion of widespread extrathoracic disease as morphology and IHC identical to DLBCL.

T lymphoblastic leukemia/lymphoma:
Use lymphoma term when confined to a mass lesion, Leukemia when there is extensive peripheral blood and bone marrow involvement. Most common in late childhood to early adulthood. Typically present acutely with symptoms related to a large mediastinal mass such as airway compromise. Mediastinal disease often centered around thymus, involving nearby lymph nodes too.

Medium-sized cells with scant cytoplasm and fine chromatin. Lots of mitoses.

IHC/Flow: (+)TdT, CD34, CD1a, CD99, CD3,

Germ Cell Tumors with associated Hematologic Malignancy:
Coexisting clonally related mediastinal germ cell tumor and a hematologic malignancy, which can be systemic or localized. Can be any type of heme malignancy, often acute leukemia. Very poor prognosis.

Metastases

Always a consideration!!

Most common = Lung.
Also consider: Breast, Esophageal, Stomach, etc...
Histiocytic and Dendritic Cell Neoplasms

Follicular Dendritic Cell Sarcoma

Intermediate-grade malignancy of follicular dendritic cells. Spindled tumor cells with indistinct cell borders, lightly eosinophilic cytoplasm, and associated lymphocytes. Oval vesicular nuclei with small nucleoli. Variable architecture. Usually only mild pleomorphism. IHC: (+) CD21, CD23, D2-40. Variable, weak CD68 & S100. Usually localized at time of Dx. Usually Adults. Rare. May arise from hyaline-vascular Castleman’s disease. Subset of patients have recurrences or metastases.

Interdigitating dendritic cell sarcoma: Rare. Very similar to FDCS (above), but derived from interdigitating dendritic cells. Plumper cells. IHC: (+) S100, (-)SOX10, CD1a, CD21, CD23, (+/-) CD68, CD45

Fibroblastic reticular cell tumor: Also similar to FDCS (above). IHC: (+) Vimentin, (-) S100, CD21, CD23. (+/-) CK, CD68

Histiocytic Sarcoma

Rare. Wide age range. Malignant proliferation of cells with histiocytic differentiation (excluding acute monocytic leukemia associated cases). Large, round, discohesive cells with abundant eosinophilic cytoplasm. Often pleomorphic. Nuclei often eccentric and vesicular.

IHC: Must express at least one histiocytic marker (e.g., CD68, CD163, or lysozyme). (-)Langerhans cell, myeloid, and follicular dendritic cell markers (in addition to epithelial and melanocytic)

Langerhans Cell Histiocytosis

Neoplastic proliferation of Langerhans cells. Discohesive cells with grooved/contorted nuclei, fine chromatin, and eosinophilic cytoplasm. Often admixed eosinophils and multinucleated giant cells.

IHC: (+)S100, CD1a, Langerin (CD207)
Molecular: Frequent BRAF V600E
Electron Microscopy: Birbeck granules
Overtly malignant cytology → Langerhans cell Sarcoma