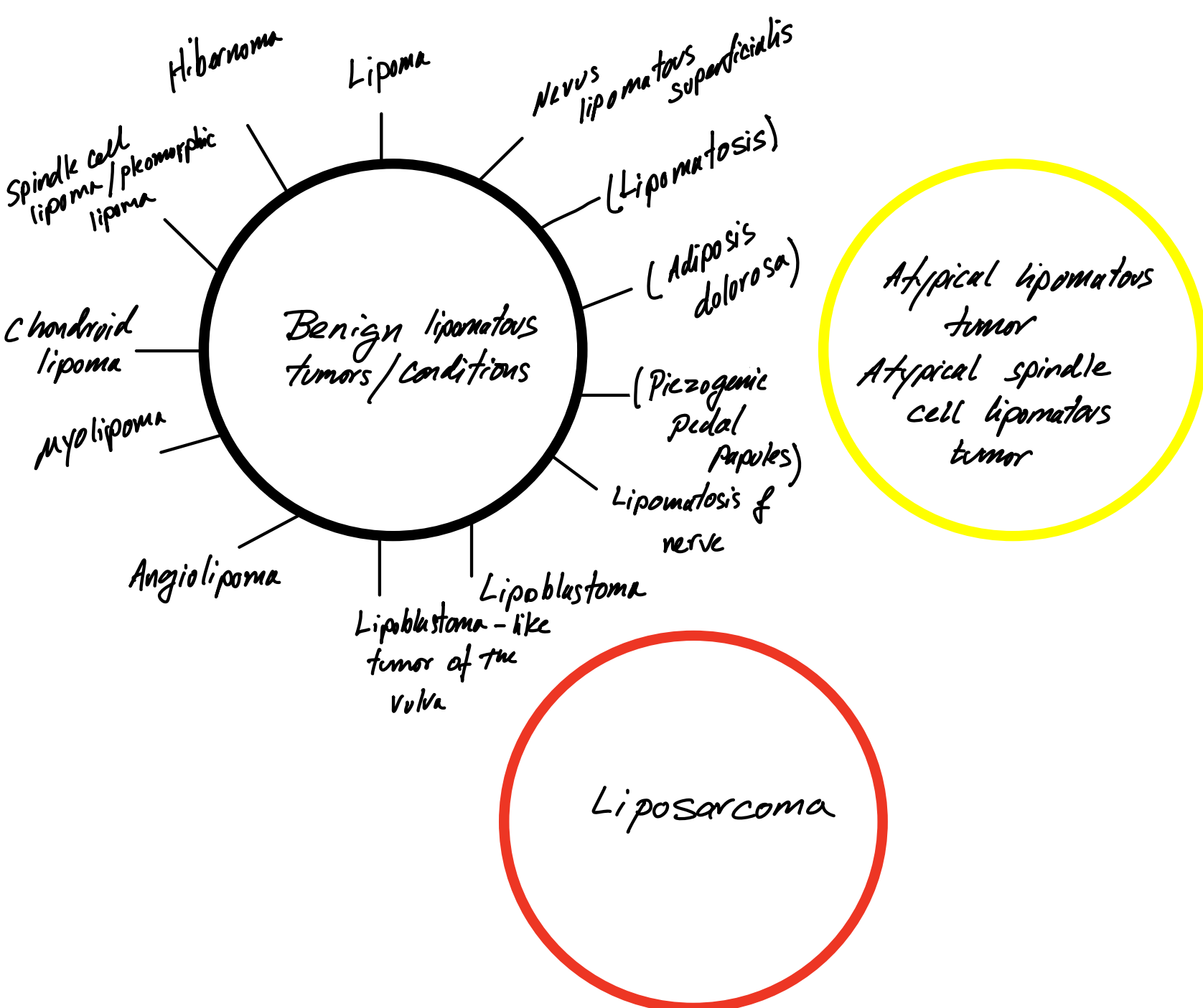
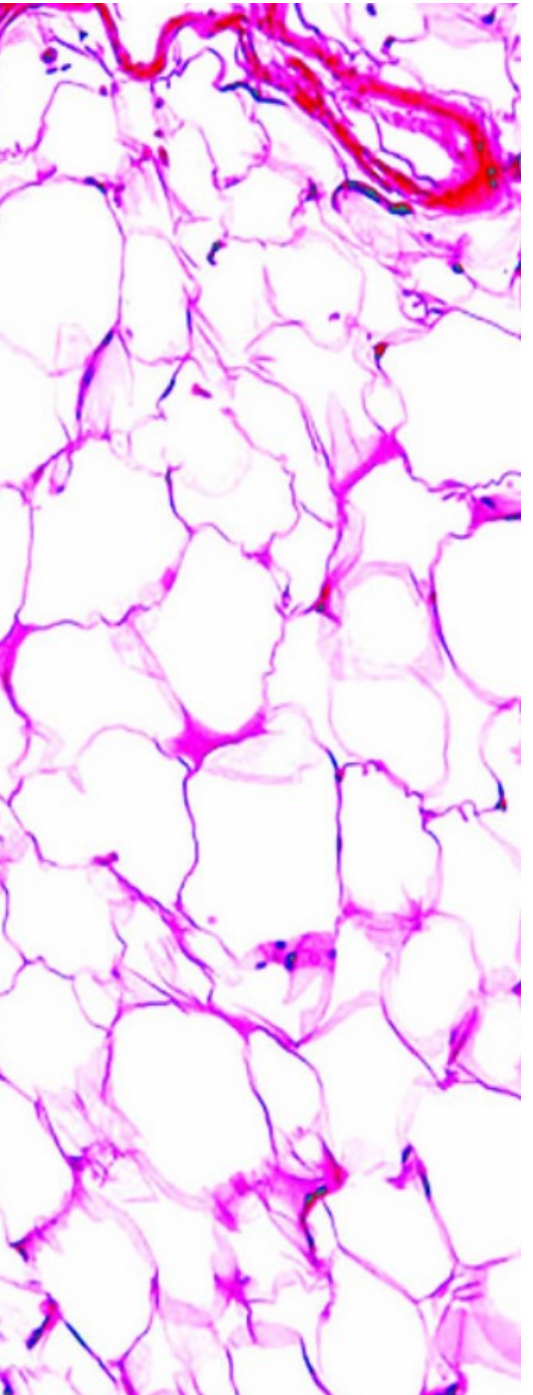


TUMORS OF FAT, MUSCLE, BONE AND CARTILAGE

Soheil S. Dadras MD-PhD

TUMORS OF ADIPOCYTES

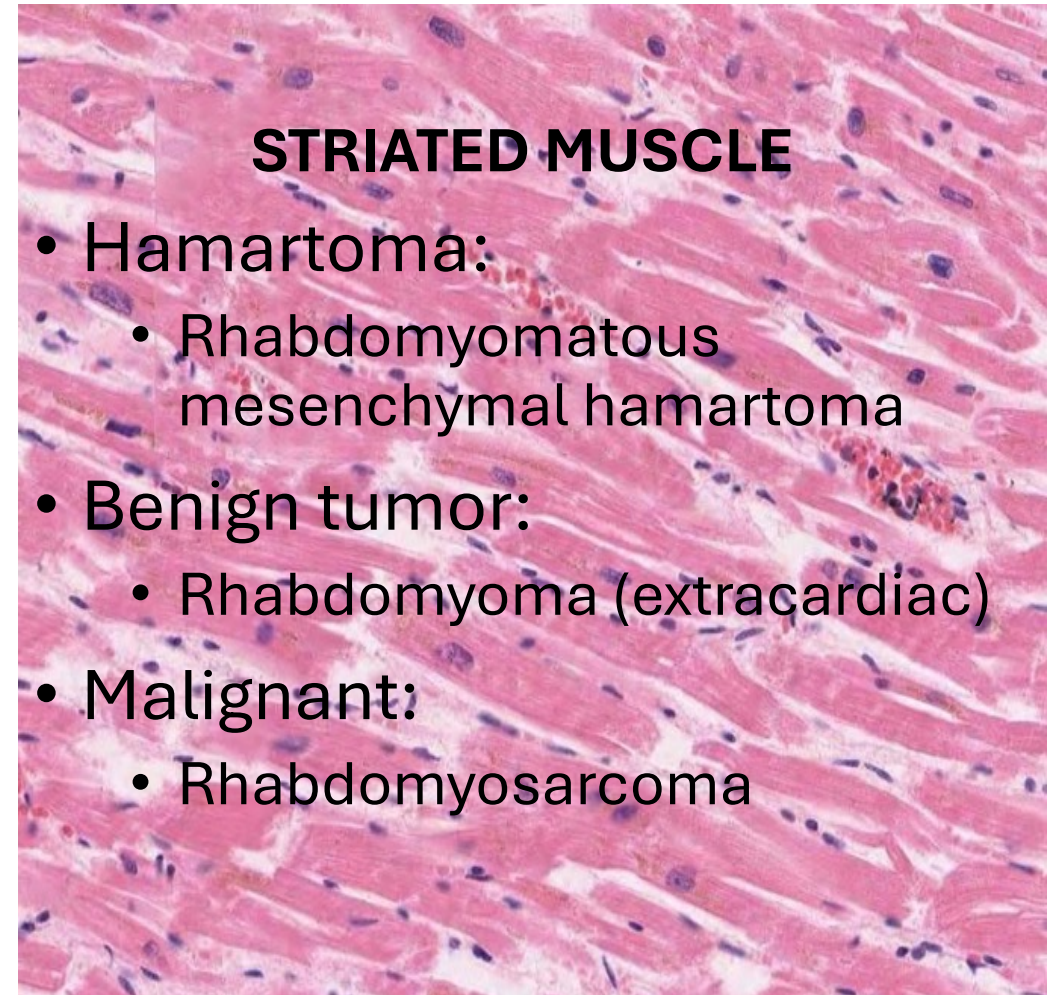


TUMORS OF MUSCLE



SMOOTH MUSCLE

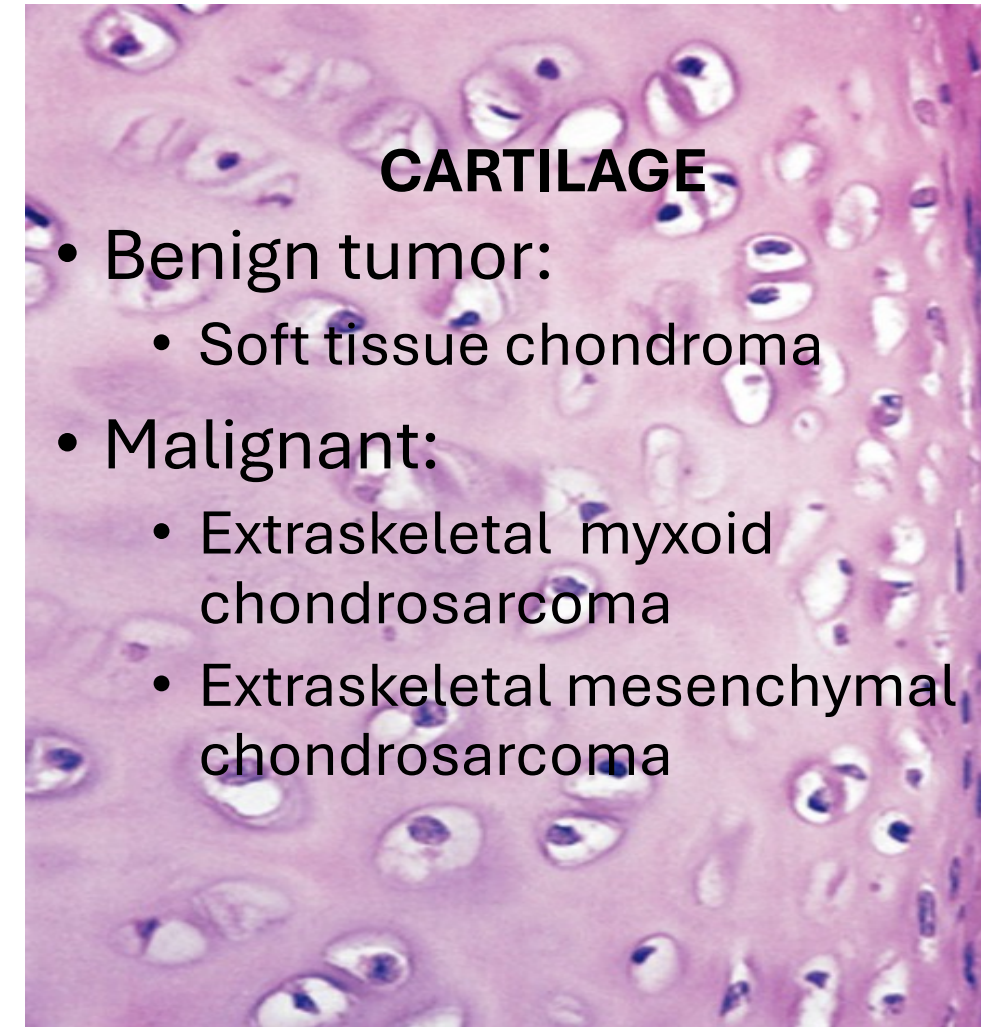
- Hamartoma:
 - Congenital smooth muscle hamartoma
- Benign tumor:
 - Pilar leiomyoma
 - Genital leiomyoma
- Malignant:
 - Leiomyosarcoma

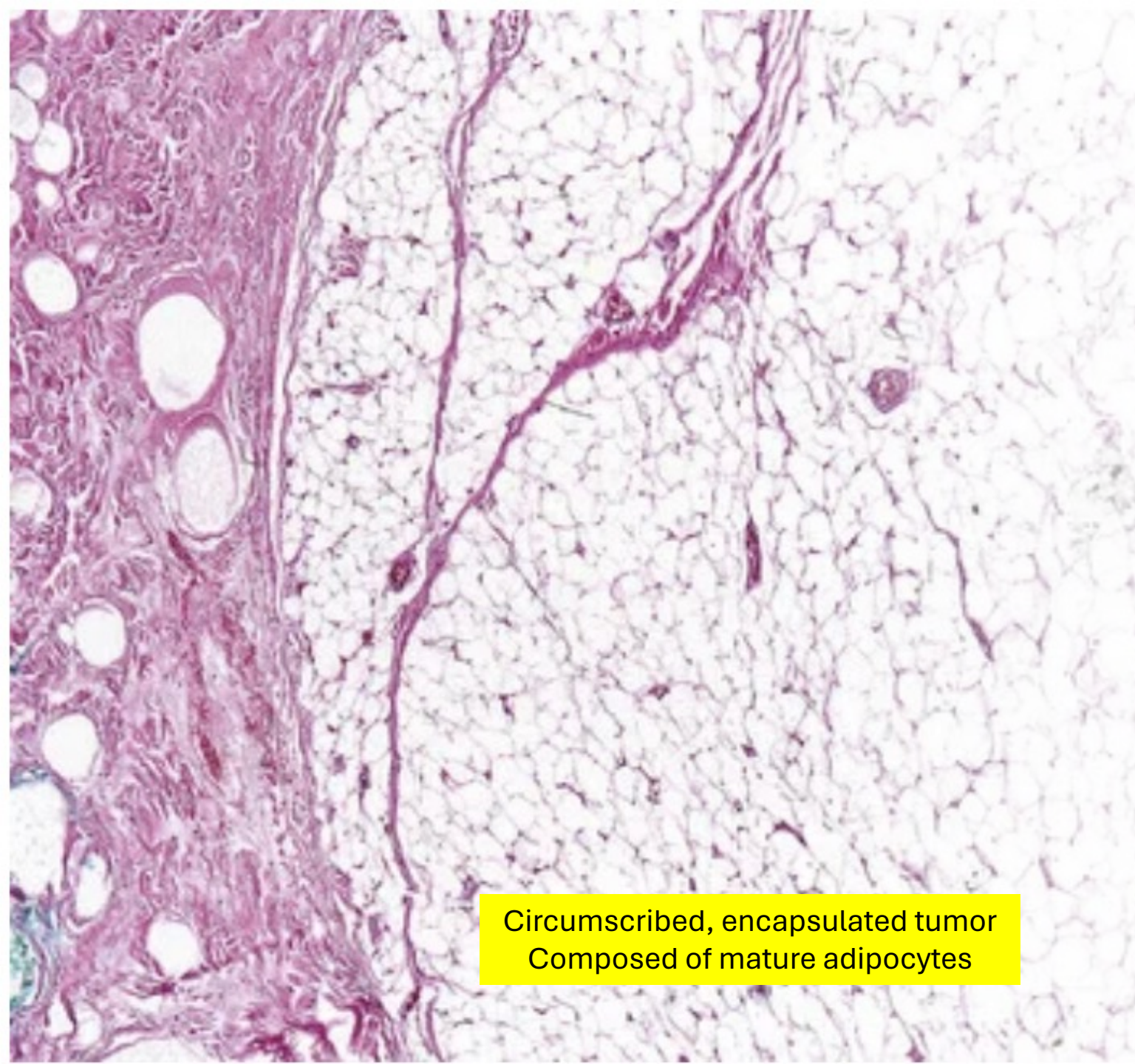


STRIATED MUSCLE

- Hamartoma:
 - Rhabdomyomatous mesenchymal hamartoma
- Benign tumor:
 - Rhabdomyoma (extracardiac)
- Malignant:
 - Rhabdomyosarcoma

TUMORS OF BONE AND CARTILAGE





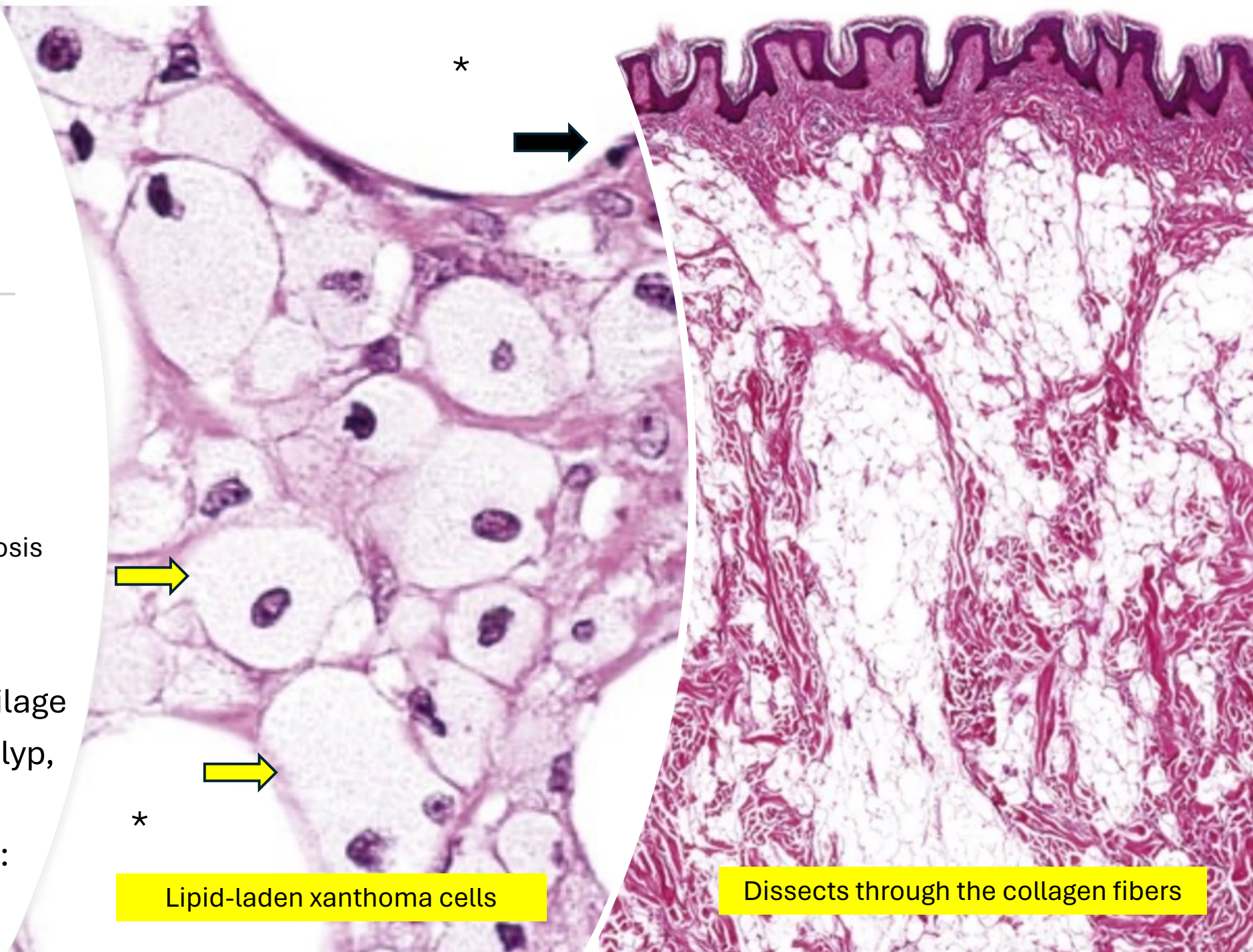
LIPOMA

- Most common CT tumors
- Uncommon in children
(Bannayan-Riley-Ruvalcaba syndrome)
- Multiple lipomas in rosiglitazone, peroxisome proliferator activator gamma agonist
- Recurrence infrequent (<5%)
- t(3;12)(q27~28;q13~15) leading to a fusion gene *HMGA2-LPP*

Circumscribed, encapsulated tumor
Composed of mature adipocytes

DERMAL LIPOMA

- Univacuolated, mature adipocytes
- Nucleus: compressed, pushed to the edge
- Degenerative changes:
 - Post-traumatic fat necrosis
 - Myxoid change
 - Fibrosis
- Other mesenchymal elements: bone or cartilage
- DDX: Fibroepithelial polyp, Nevus lipomatosus superficialis, Atypical lipomatous tumor (IHC: MDM2+, CDK4+)





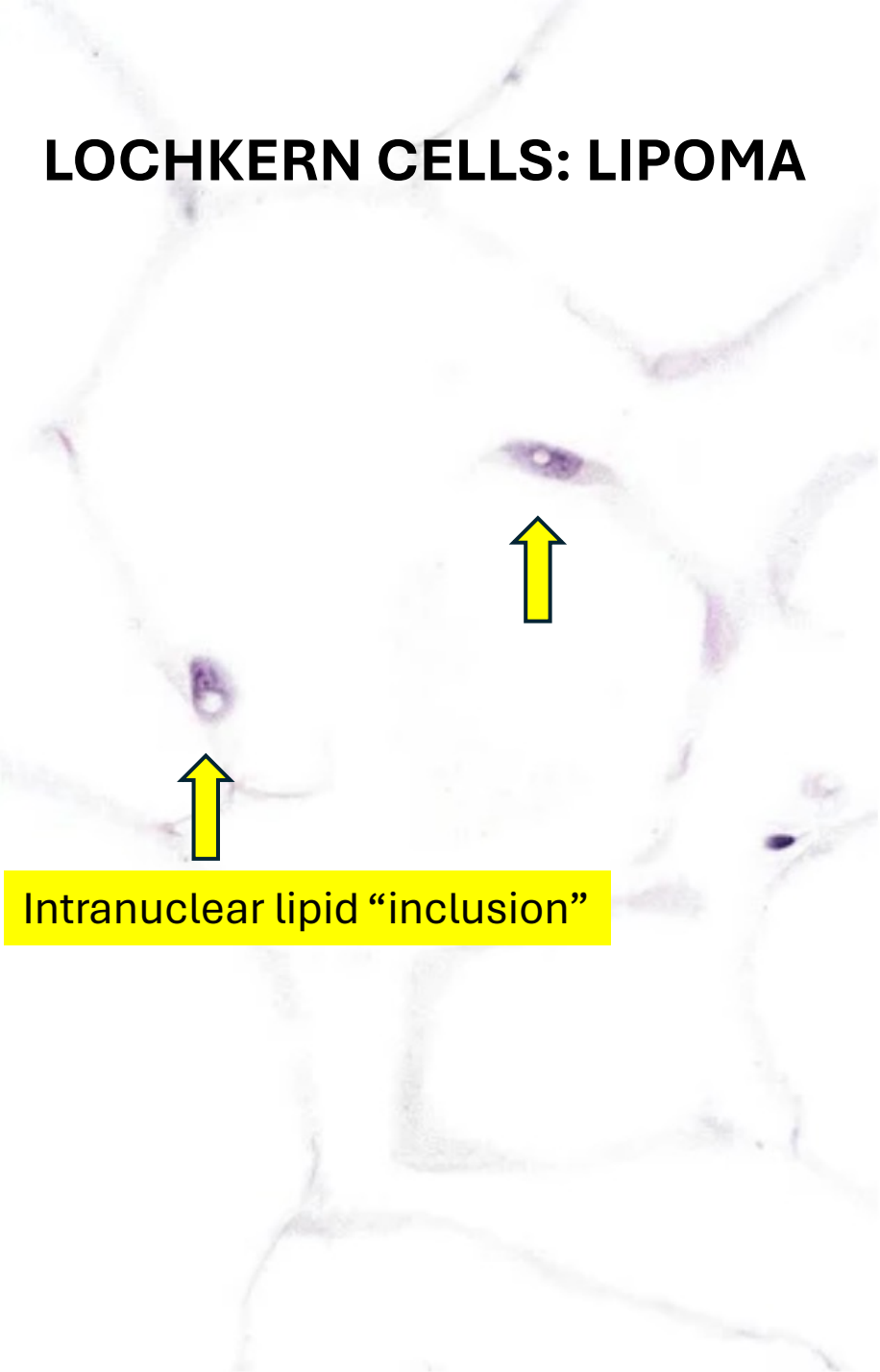
Mature
adipocytes

↑
Fibrosis

Myxoid change

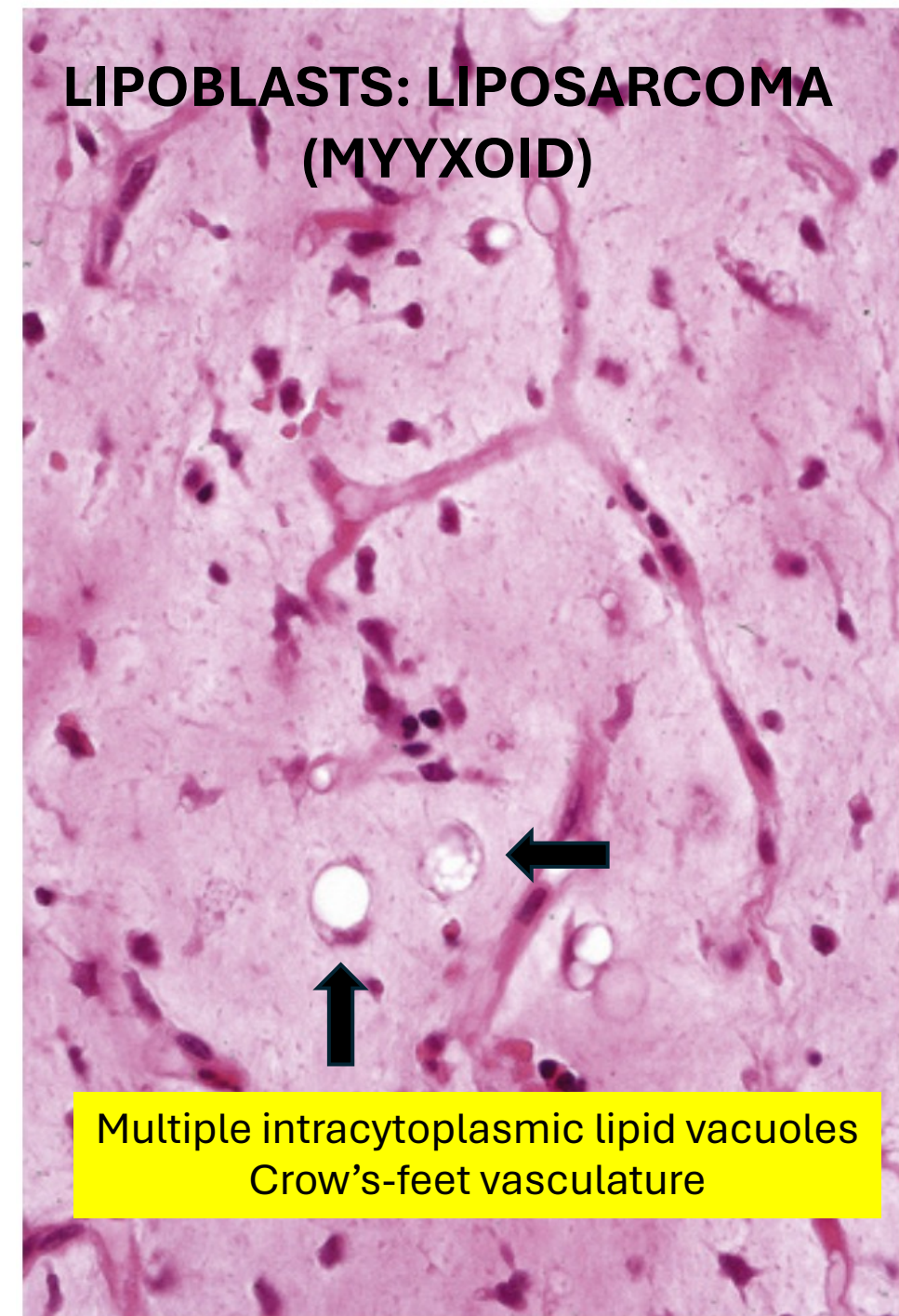
Myxofibrolipoma

LOCHKERN CELLS: LIPOMA



Intranuclear lipid “inclusion”

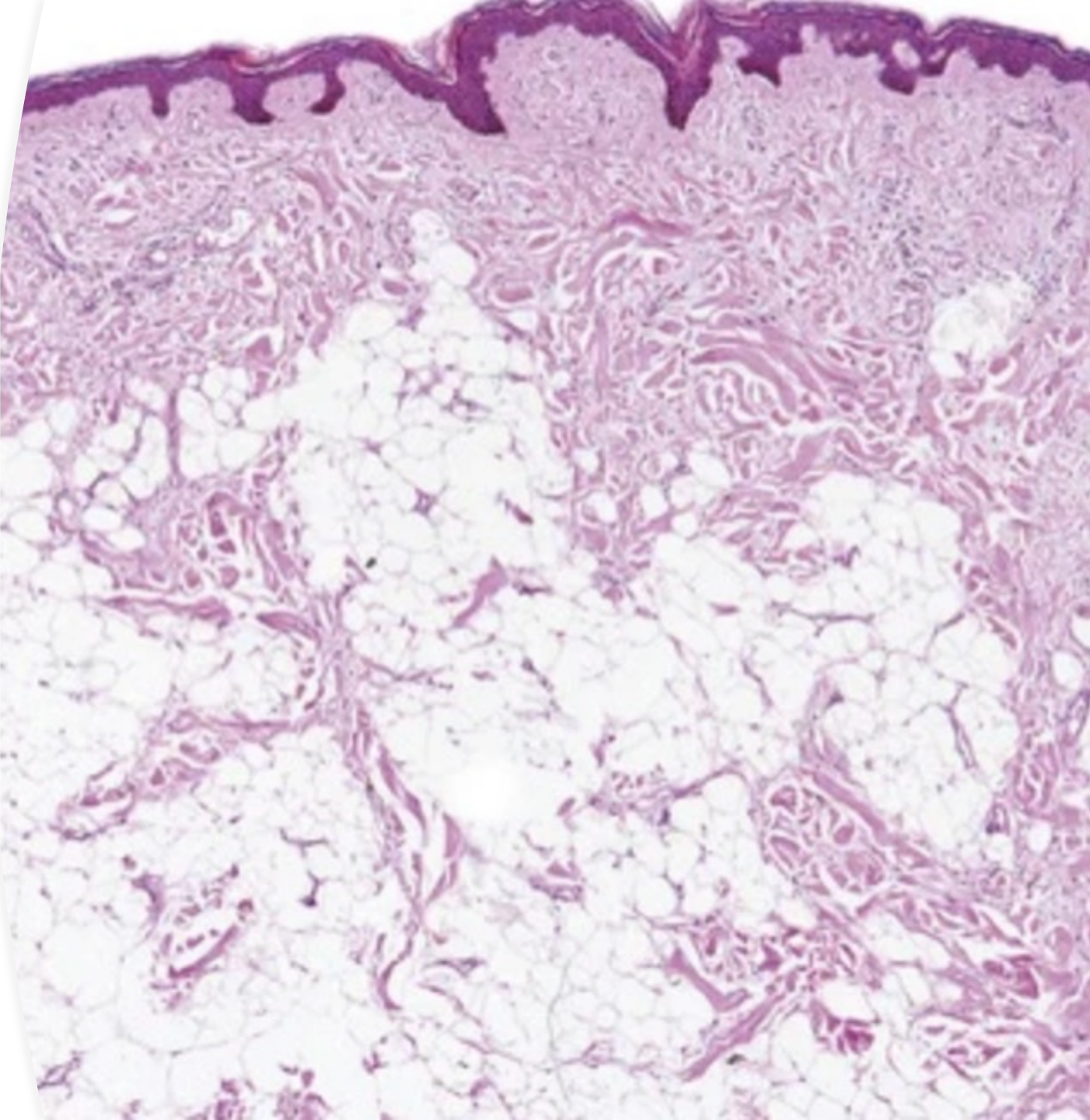
LIPOBLASTS: LIPOSARCOMA (MYXOID)



Multiple intracytoplasmic lipid vacuoles
Crow's-feet vasculature

NEVUS LIPOMATOSUS SUPERFICIALIS

- Uncommon connective tissue nevus
- Multiple papular, polypoid or plaques
- Buttocks, upper thigh or lower back



LIPOMATOSIS, ADIPOSIS DOLOROSA AND PIEZOGENIC PEDAL PAPULES

- HISTOLOGY:
 - Normal, unencapsulated adipose tissue herniating into the dermis
 - DDX: dermal lipoma
- CLINICAL:
 - Lipomatosis: symmetric/diffuse or asymmetric
 - Syndromic association
 - Etiology: HIV lipodystrophy, exogenous/endogenous production of steroids, or idiopathic
 - Adiposis dolorosa : increased fat in painful, plaque-like distribution
 - Piezogenic pedal papules: multiple skin-colored papules on the heels

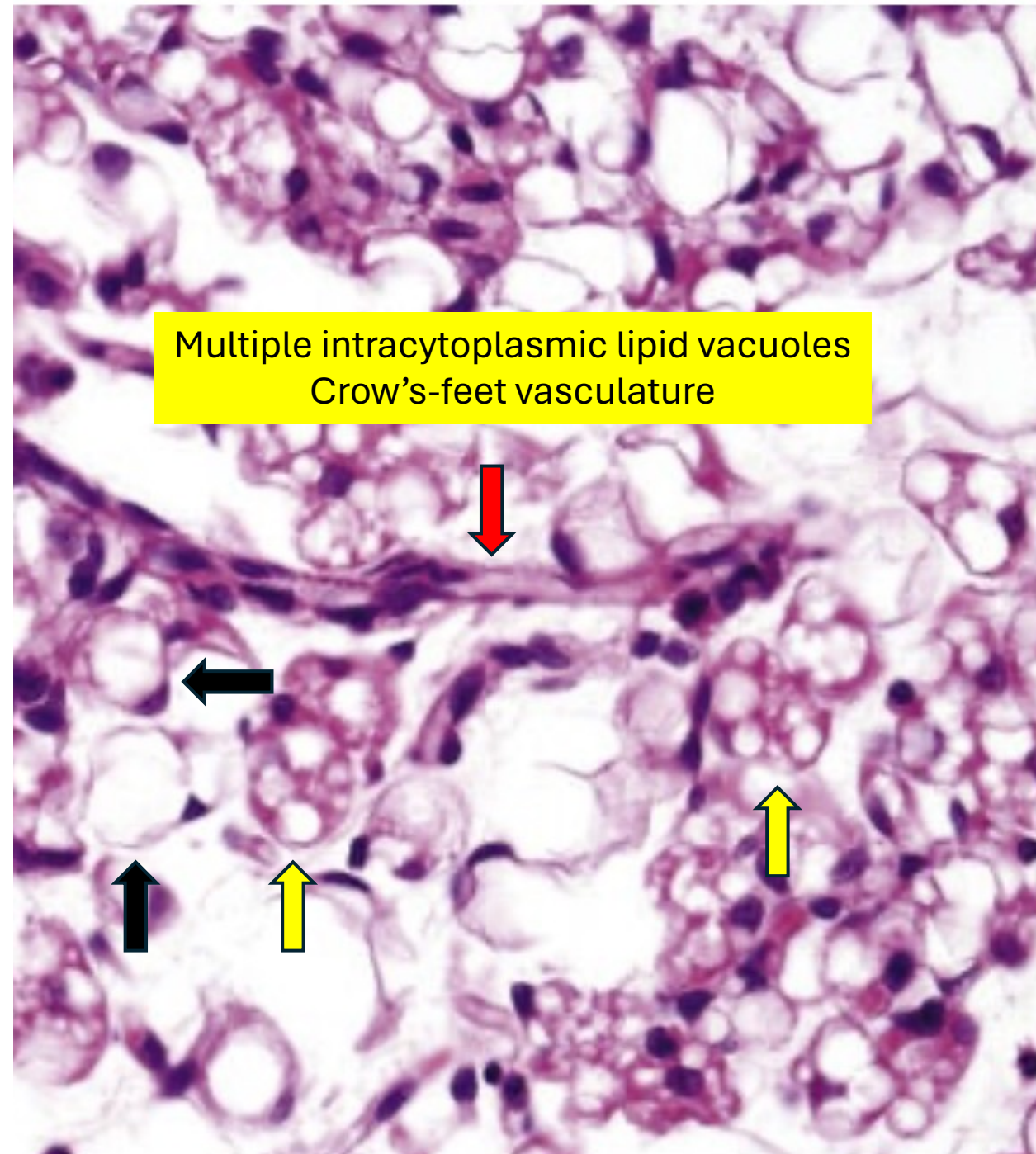
LIPOMATOSIS OF NERVE

- Hamartoma of nerve as well as adipose and fibrous tissue
- Within epineurium of nerve
- Concentric perineural fibrosis
- Rare bone formation

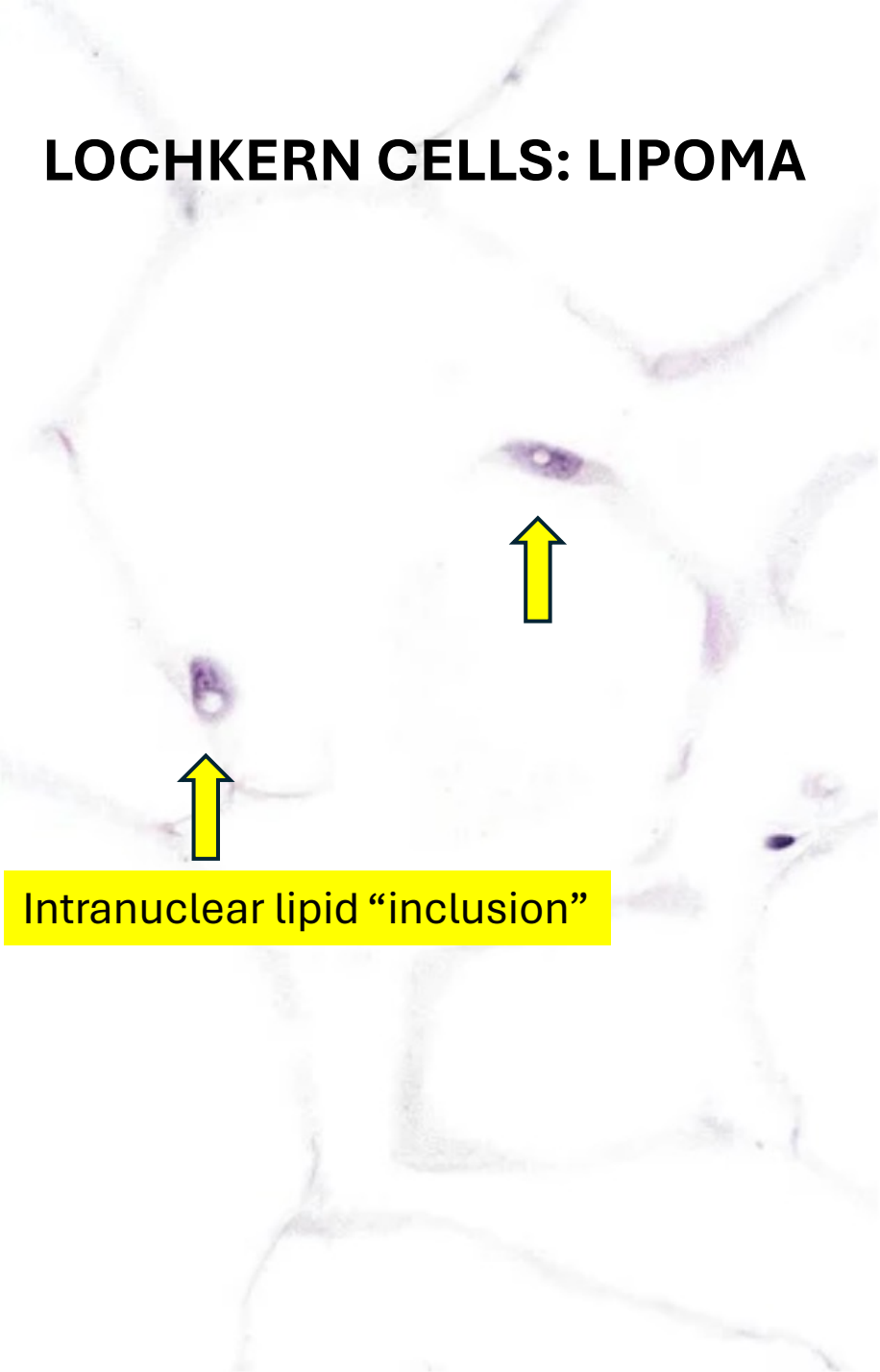


LIPOBLASTOMA

- Counterpart of lipoma in infancy/childhood
- Resembles hemangioma
- Trunk > extremities > head and neck
- Rearrangement of 8q11~q13
 - Overexpression of *PLAG1* oncogene
- Recapitulates developing fat:
 - Mature adipocytes, lipoblasts and perilipoblasts
- IHC: S100+, CD34+, PLAG1+, rare p16+

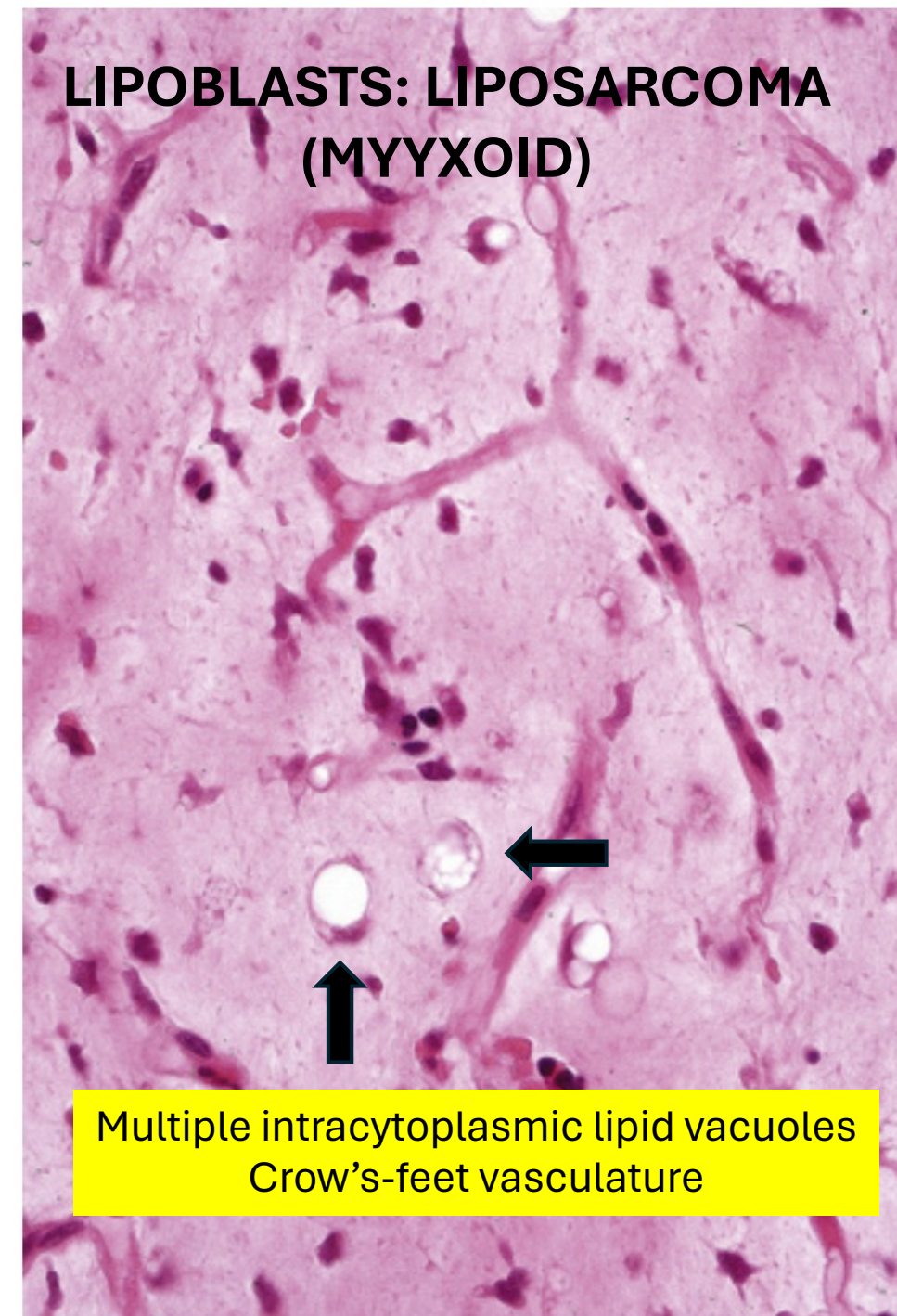


LOCHKERN CELLS: LIPOMA



Intranuclear lipid "inclusion"

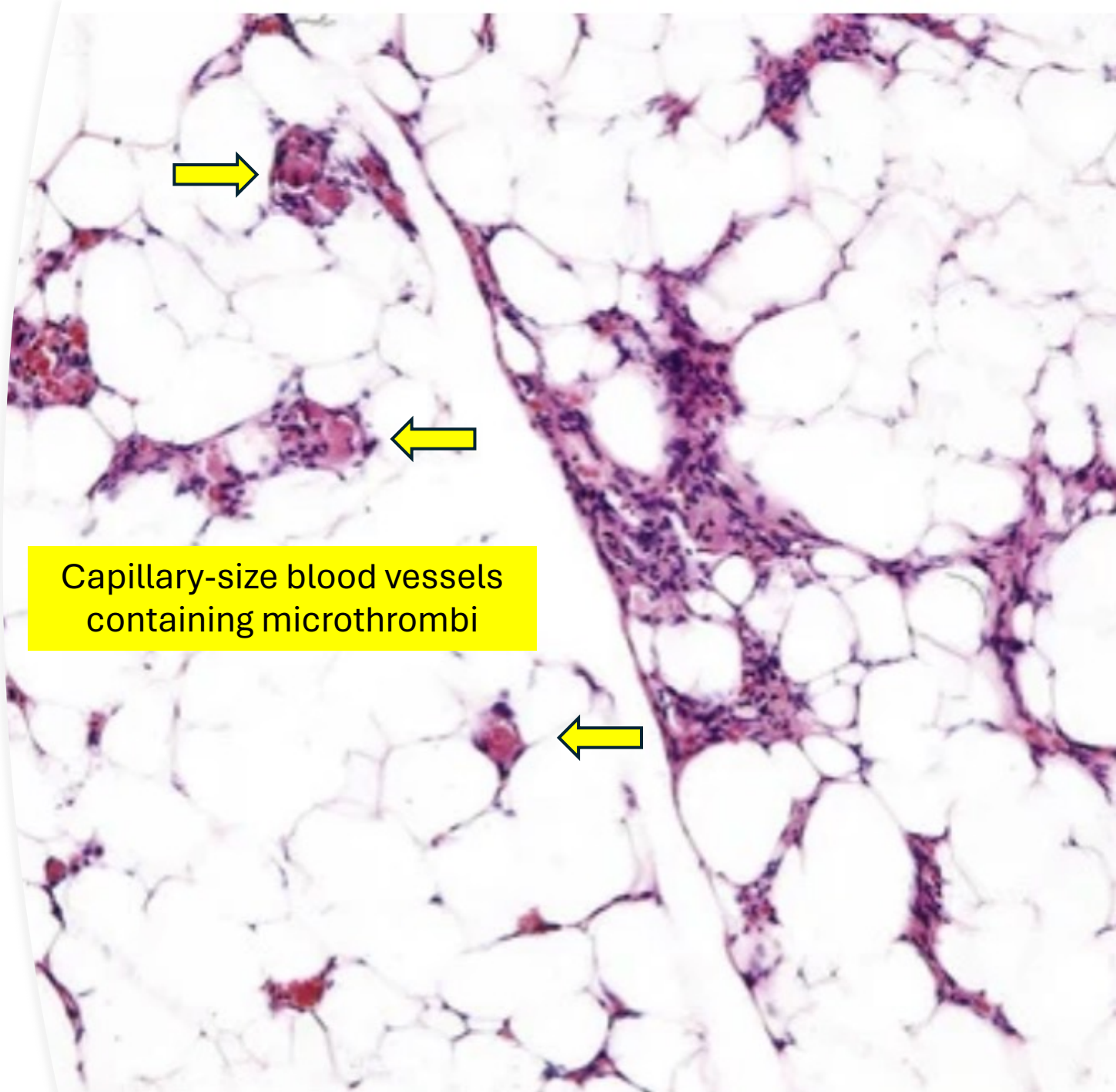
LIPOBLASTS: LIPOSARCOMA (MYXOID)

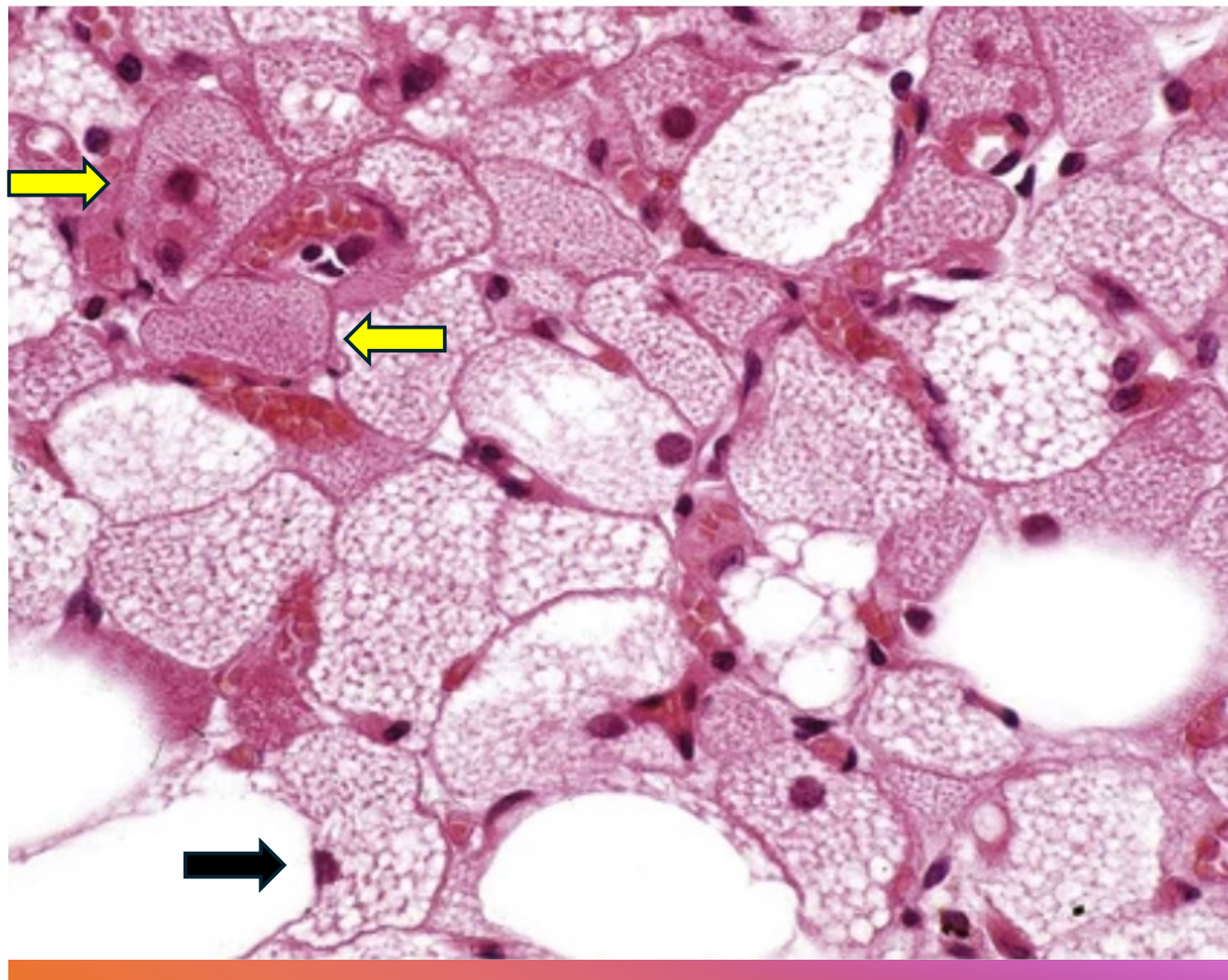


Multiple intracytoplasmic lipid vacuoles
Crow's-feet vasculature

ANGIOLIPOMA

- Young adults, the subcutis of forearm and trunk
- Tender, painful, red-blue discoloration
- Multiple indinavir and saquinavir
- Aberrant structure of chromosome 13, expression of protein kinase D2 and HMGA2
- DDX: capillary hemangioma and KHE





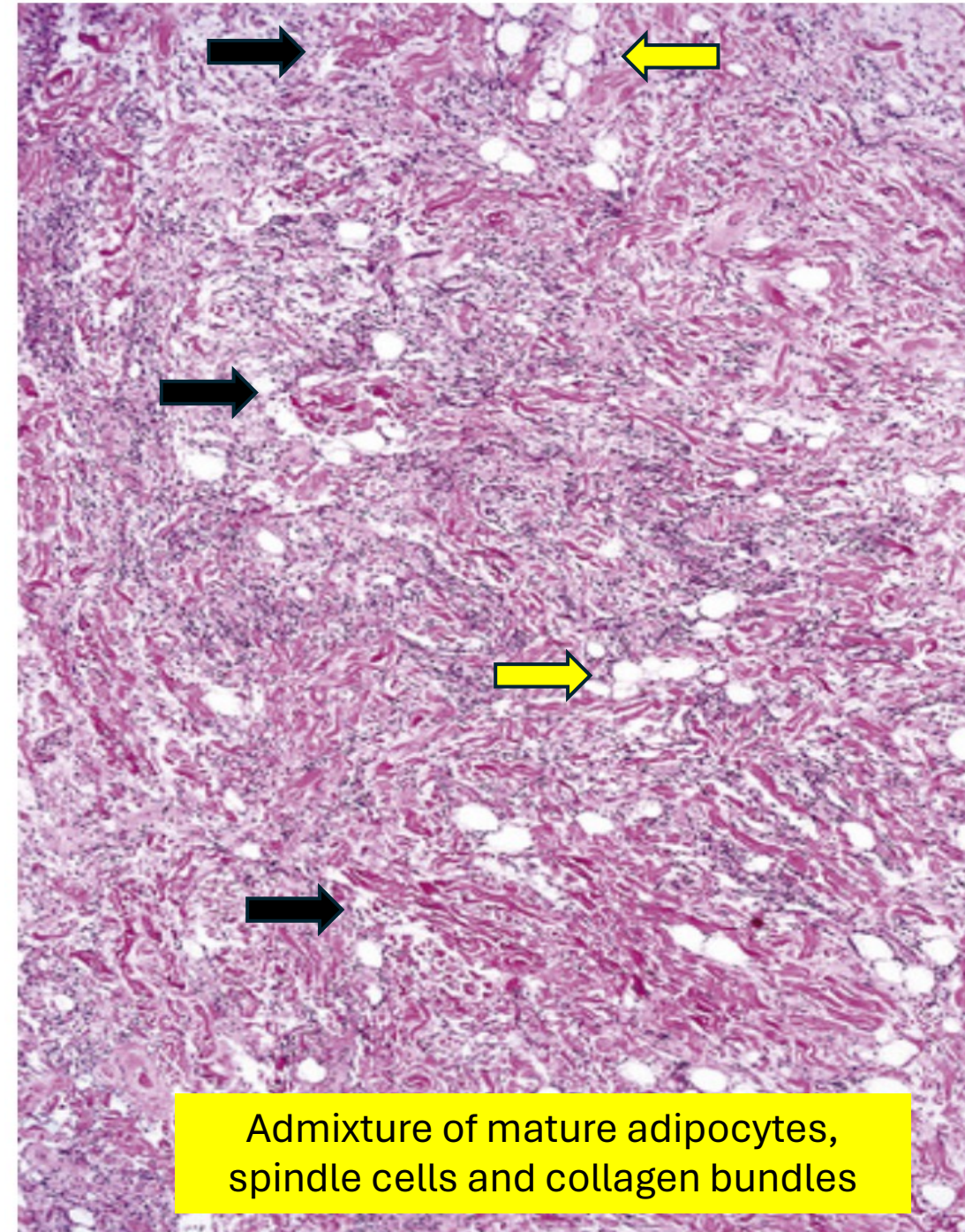
Mixed mature adipocytes and multivacuolated adipocytes with granular eosinophilic cytoplasm

HIBERNOMA

- Rare, benign brown fat in young adults
- Interscapular space, axillae, chest wall and head and neck
- Rearrangement of 11q13
- *MEN1* and/or *AIP* loss
- Encapsulated, lobulated mass
- ± Lipoblast-like cells
- IHC: MDM2-, CDK4-

SPINDLE CELL LIPOMA/ PLEOMORPHIC LIPOMA

- Uncommon variants of lipoma (benign)
- Histologically concerning to the novice
- Posterior neck, shoulder or upper back of adult males
- Aberrant structure of chromosomes 13 and 16
- DDX: liposarcoma, soft tissue neoplasm NOS
- IHC: spindle cells CD34+, S100-

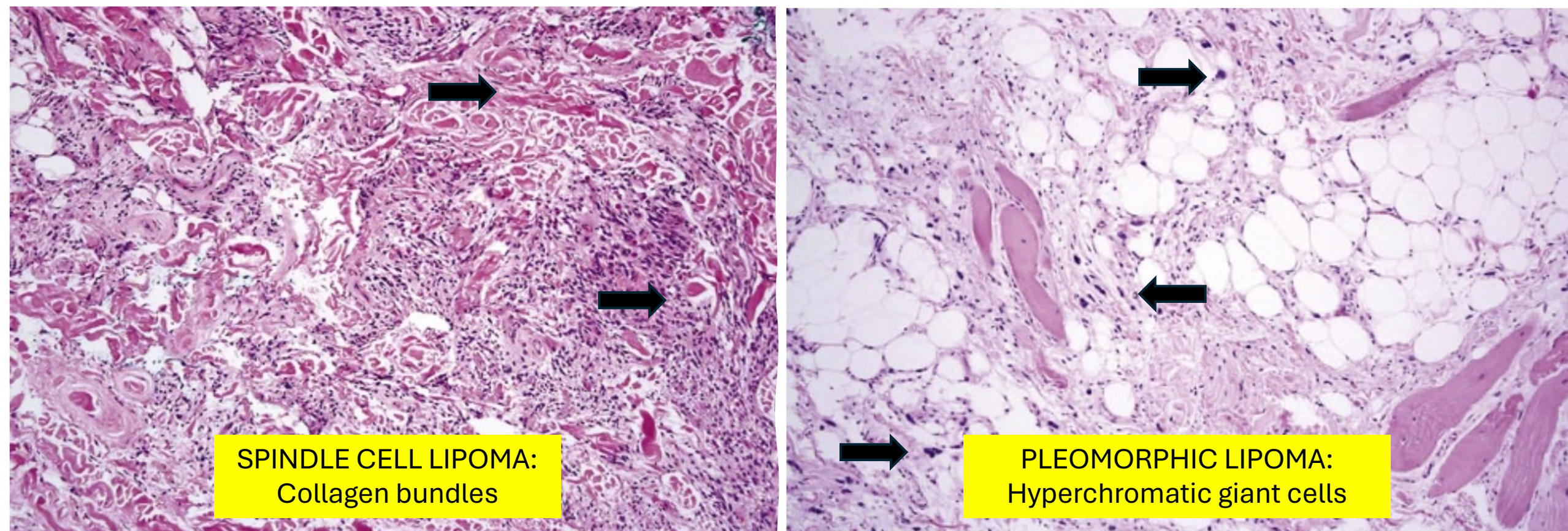


Morphologic continuum

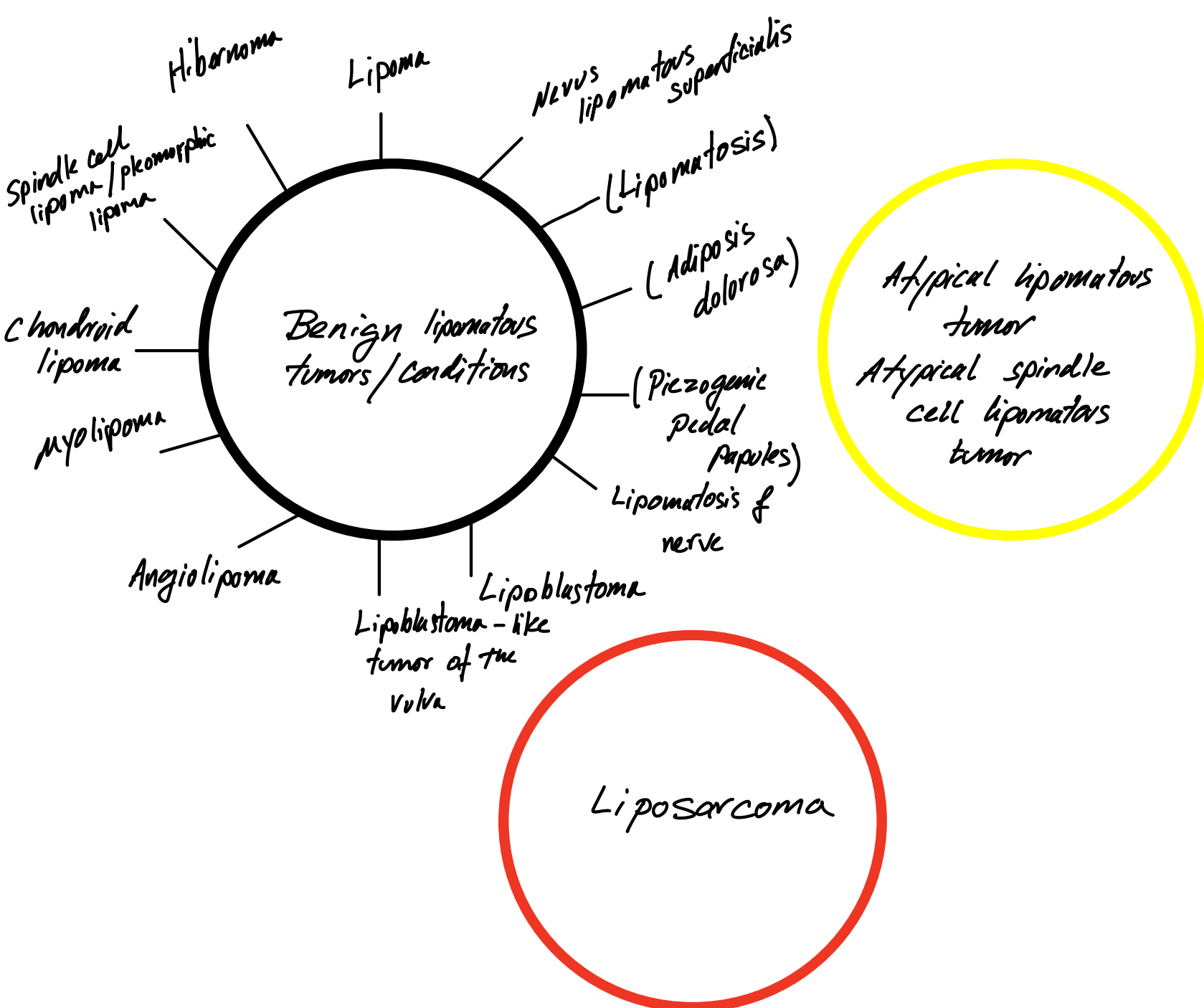
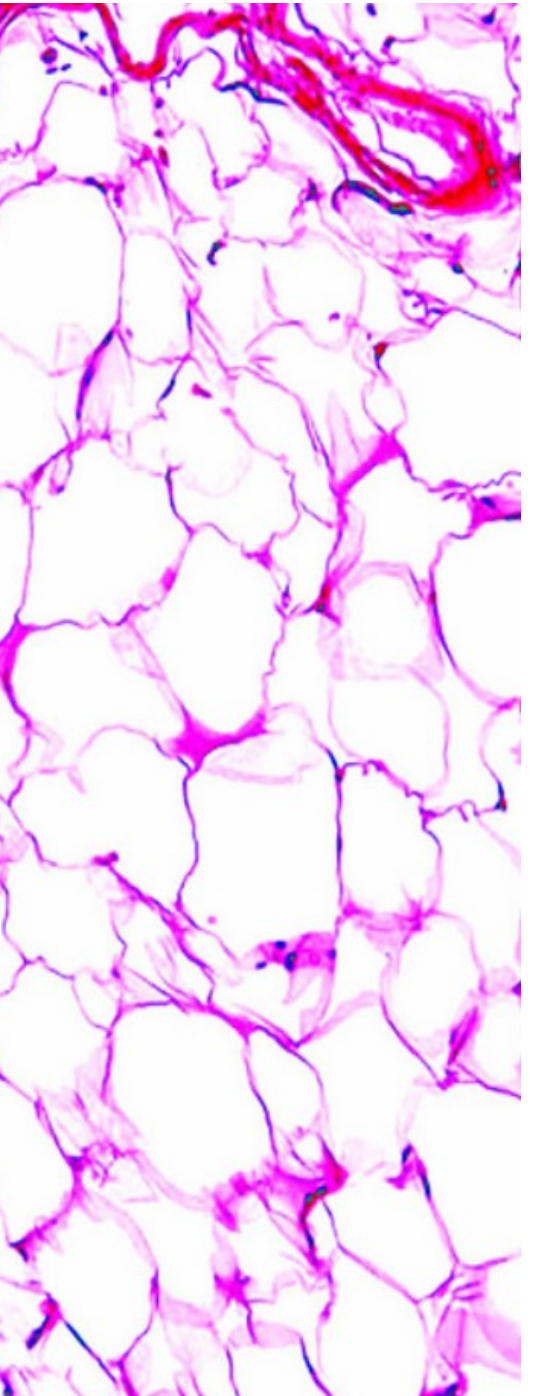
SPINDLE CELL LIPOMA:
Collagen bundles

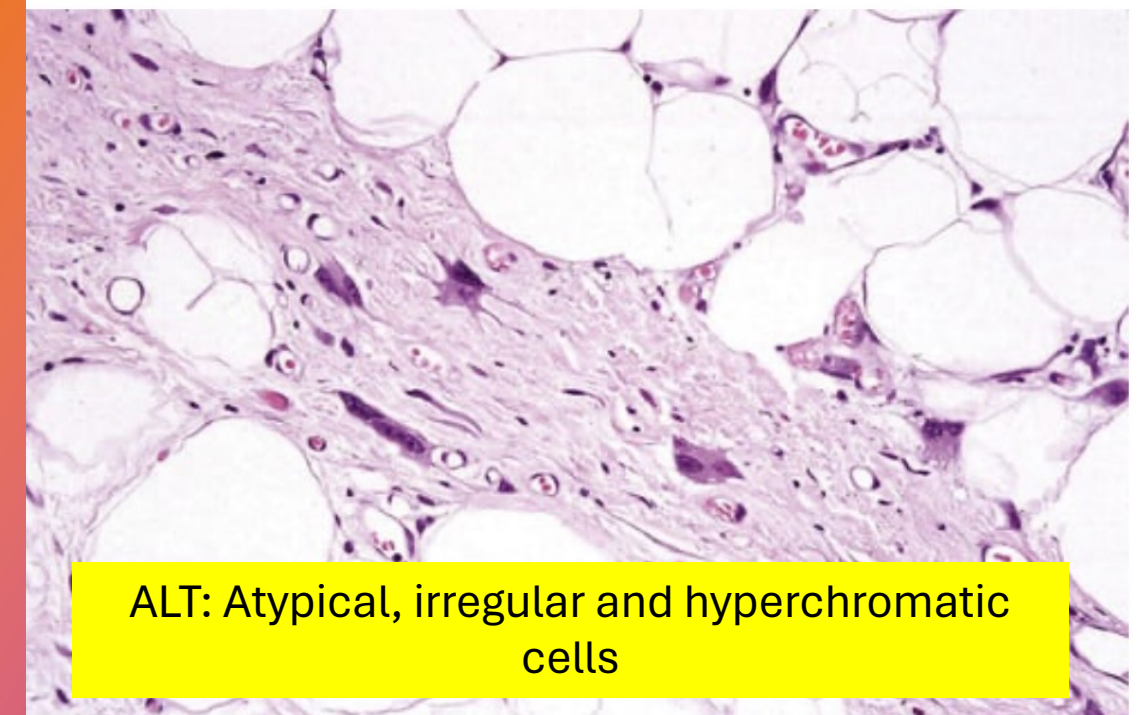
PLEOMORPHIC LIPOMA:
Hyperchromatic giant cells

- Ill-defined subcutaneous mass
- Univacuolated adipocytes and slender spindled cells
- Collagen bundles
- Myxoid degeneration (pseudovascular spaces)
- Fat and vascular components may vary
- Mast cells

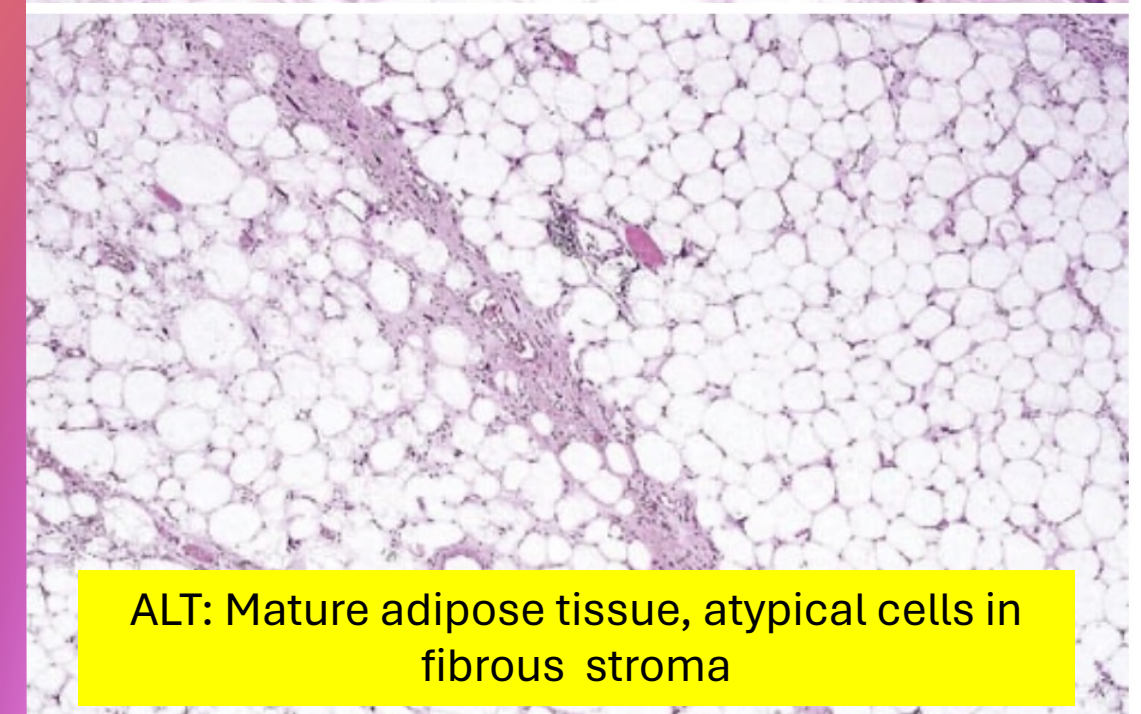


TUMORS OF ADIPOCYTES





ALT: Atypical, irregular and hyperchromatic cells

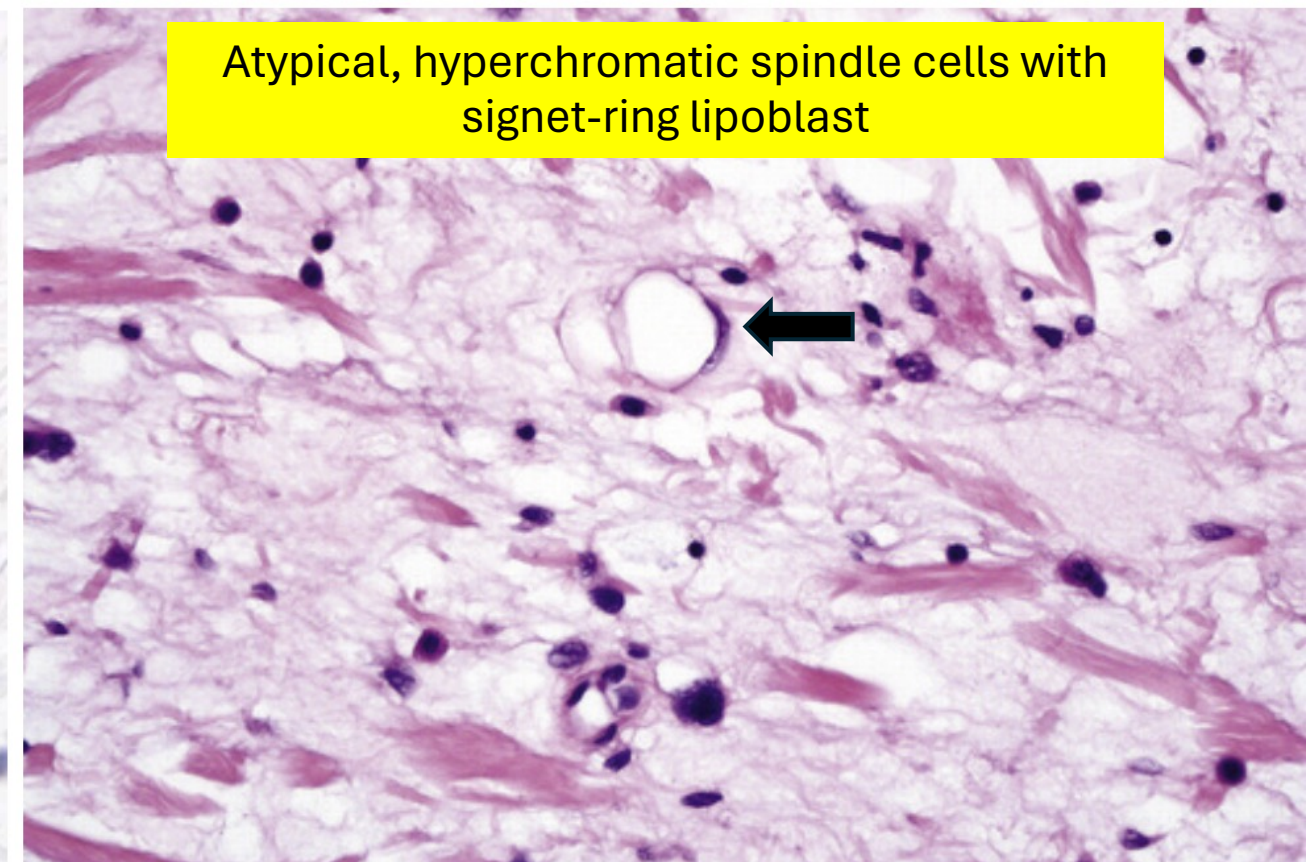
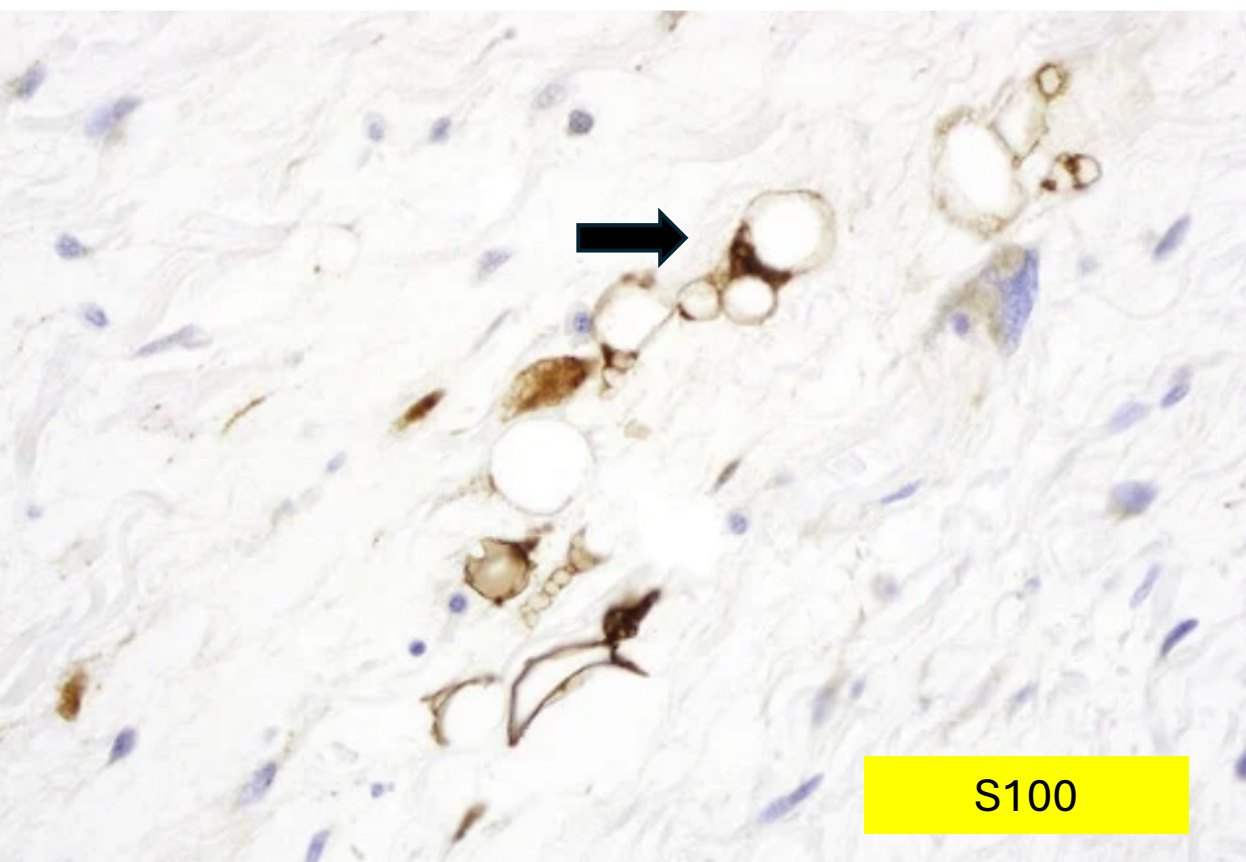


ALT: Mature adipose tissue, atypical cells in fibrous stroma

ADIPOCYTIC TUMORS OF INTERMEDIATE MALIGNANCY

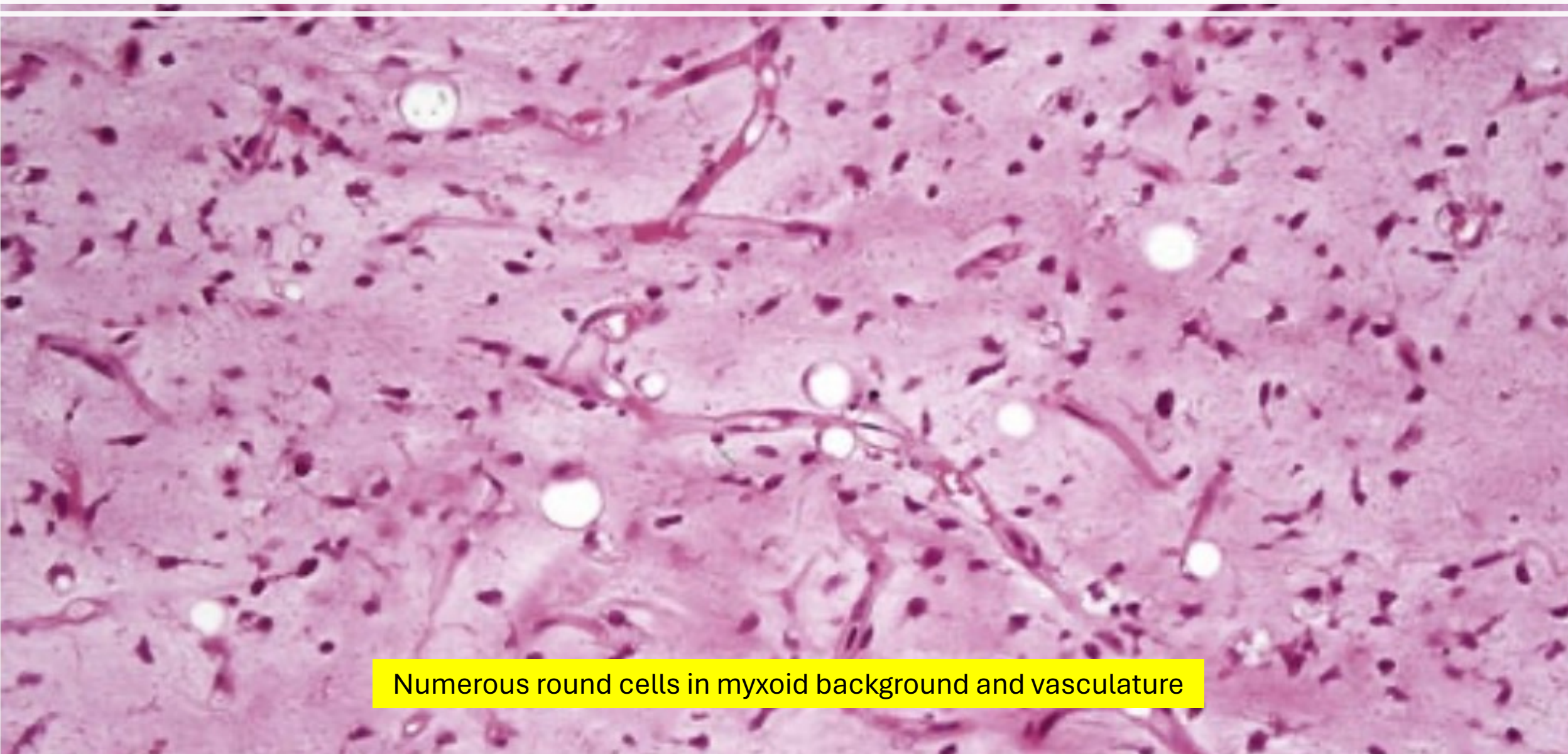
- Locally aggressive, no metastatic potential
 - Atypical lipomatous tumor (ALT)
 - Atypical spindle cell lipomatous tumor
- Local recurrence, if incompletely excised
- Deep seated: subcutis, skeletal muscle, retroperitoneum, mediastinum, and spermatic cord
- Older males, trunk
- Ring 12q13~15, amplified *MDM2* and *CDK4* (IHC+)
- DDX: Lipoma (lochkorn cells)

ATYPICAL SPINCEL CELL LIPOMATOUS TUMOR



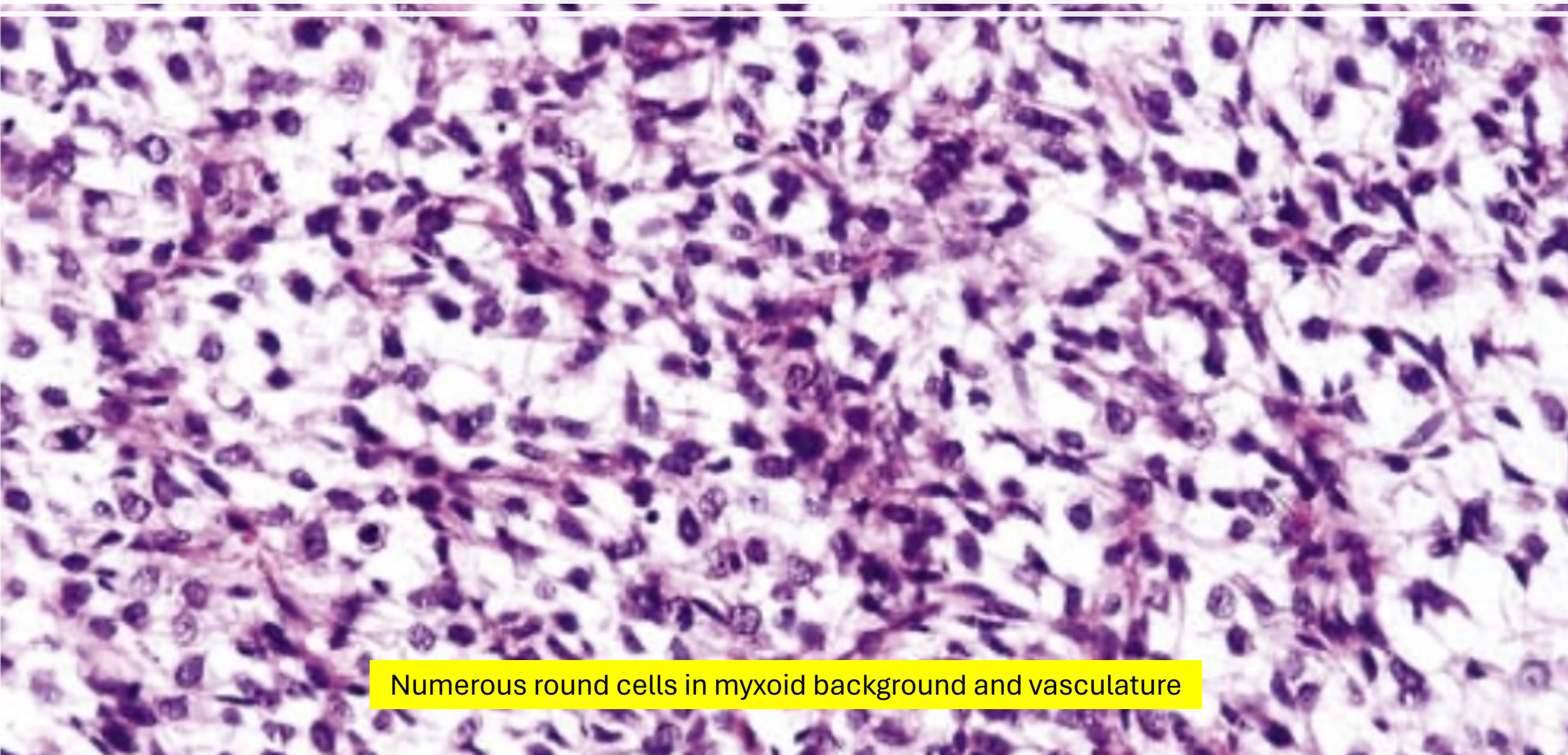
- Middle-aged males: hands, feet and limb girdles
- Local recurrence: 12%, no metastatic potential
- Heterozygous deletion of *RB1* (loss of expression by IHC)
- IHC: *RB1*-, *CD34*+, *S100*+, *MDM2* ±, *CDK4*±
- DDX: pleomorphic lipoma, DFSP, low-grade malignant peripheral nerve sheath tumor, low-grade dedifferentiated liposarcoma

MYXOID LIPOSARCOMA



Numerous round cells in myxoid background and vasculature

MYXOID LIPOSARCOMA



Numerous round cells in myxoid background and vasculature

PLEOMORPHIC LIPOSARCOMA



Lipoblasts in pleomorphic cellular background

LIPOSARCOMA VARIANTS

- ALT/Dedifferentiated/Well-differentiated
 - Abrupt transition to high-grade, nonlipogenic sarcoma
 - IHC: MDM2+, CDK4+
- Myxoid
 - Middle-aged adults, rare in children
 - Lower limb, thigh
 - ‘Crows-feet’ pattern small thin-walled capillaries
 - Local recurrence
 - 30% metastasize
 - Poor prognosis: p53 overexpression, necrosis
- Pleomorphic
 - Elderly, deep-seated, limbs
 - Rapid growth, high local recurrence and metastasis
- DDX: atypical lipomatous tumor, other soft tissue neoplasms (fat-free) myxofibrosarcoma

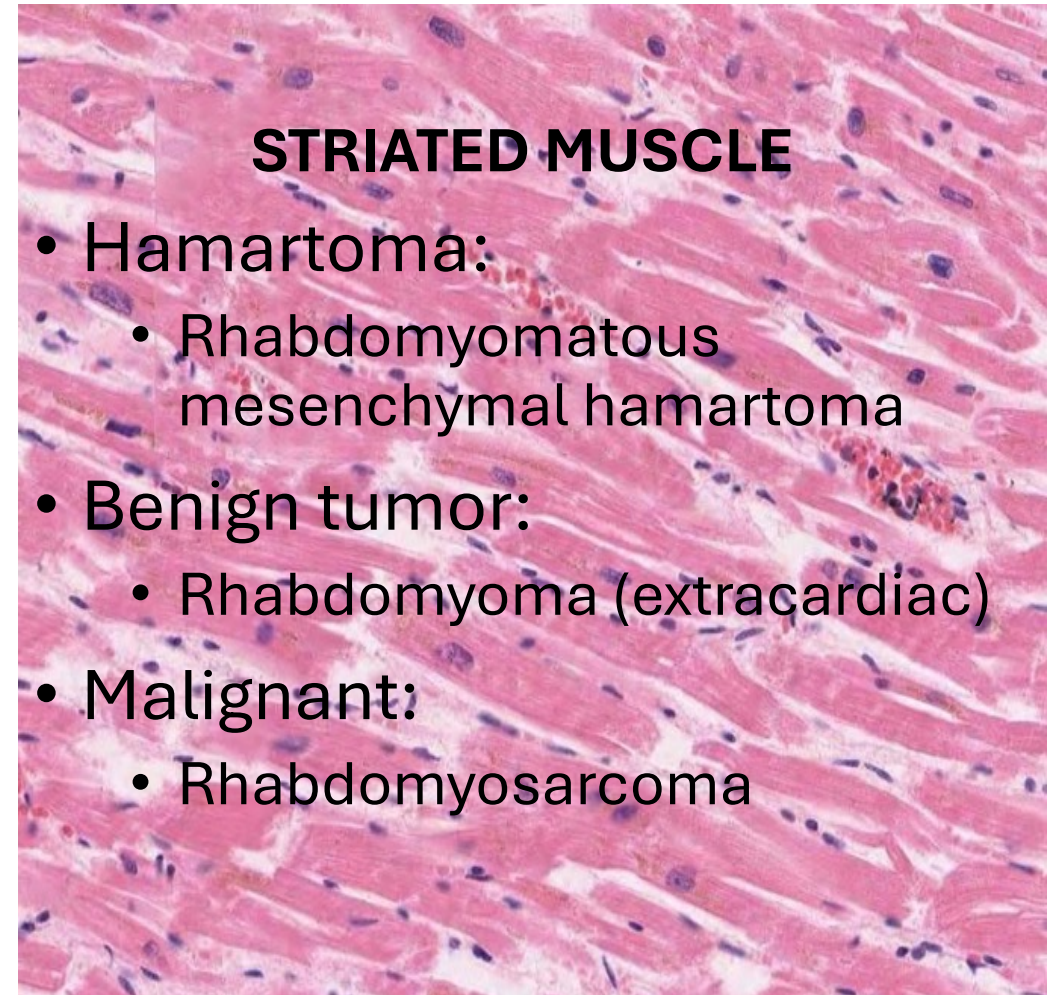


TUMORS OF MUSCLE



SMOOTH MUSCLE

- Hamartoma:
 - Congenital smooth muscle hamartoma
- Benign tumor:
 - Pilar leiomyoma
 - Genital leiomyoma
- Malignant:
 - Leiomyosarcoma



STRIATED MUSCLE

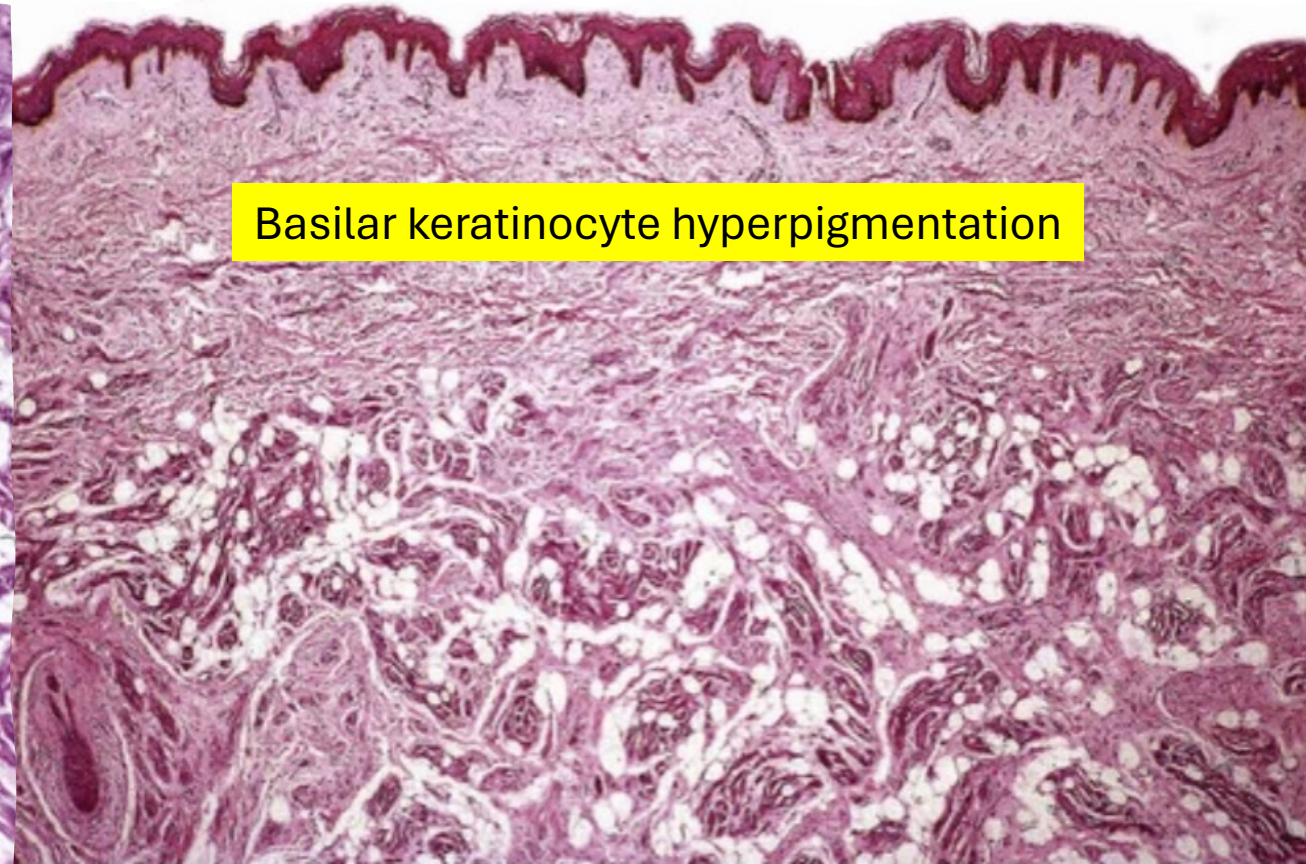
- Hamartoma:
 - Rhabdomyomatous mesenchymal hamartoma
- Benign tumor:
 - Rhabdomyoma (extracardiac)
- Malignant:
 - Rhabdomyosarcoma

CONGENITAL SMOOTH MUSCLE HAMARTOMA

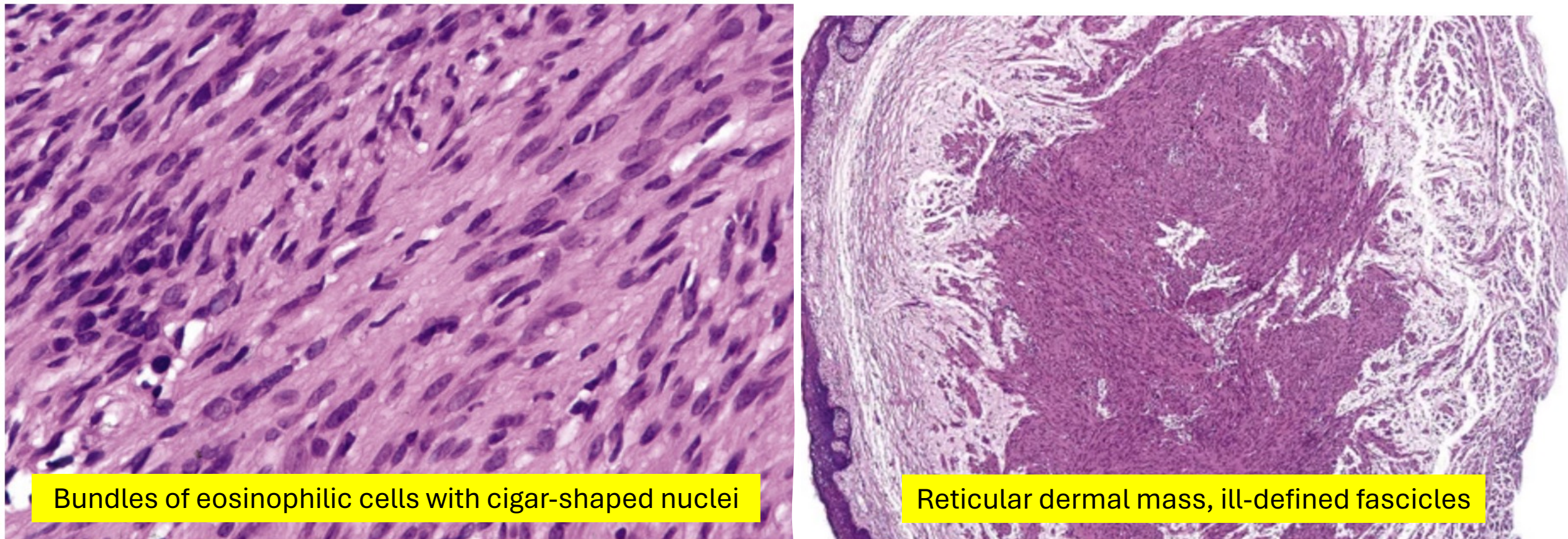
- Male infants
- Lumbosacral, proximal thighs
- Indurated, hyperpigmented macule/plaque with coarse hairs
- DDX: Becker nevus (acquired, hypertrichosis, subtle histology)
- IHC: SMA+, desmin+, and h-caldesmon+ (smooth muscle)



Bundles of smooth muscle and adipocytes



Basilar keratinocyte hyperpigmentation



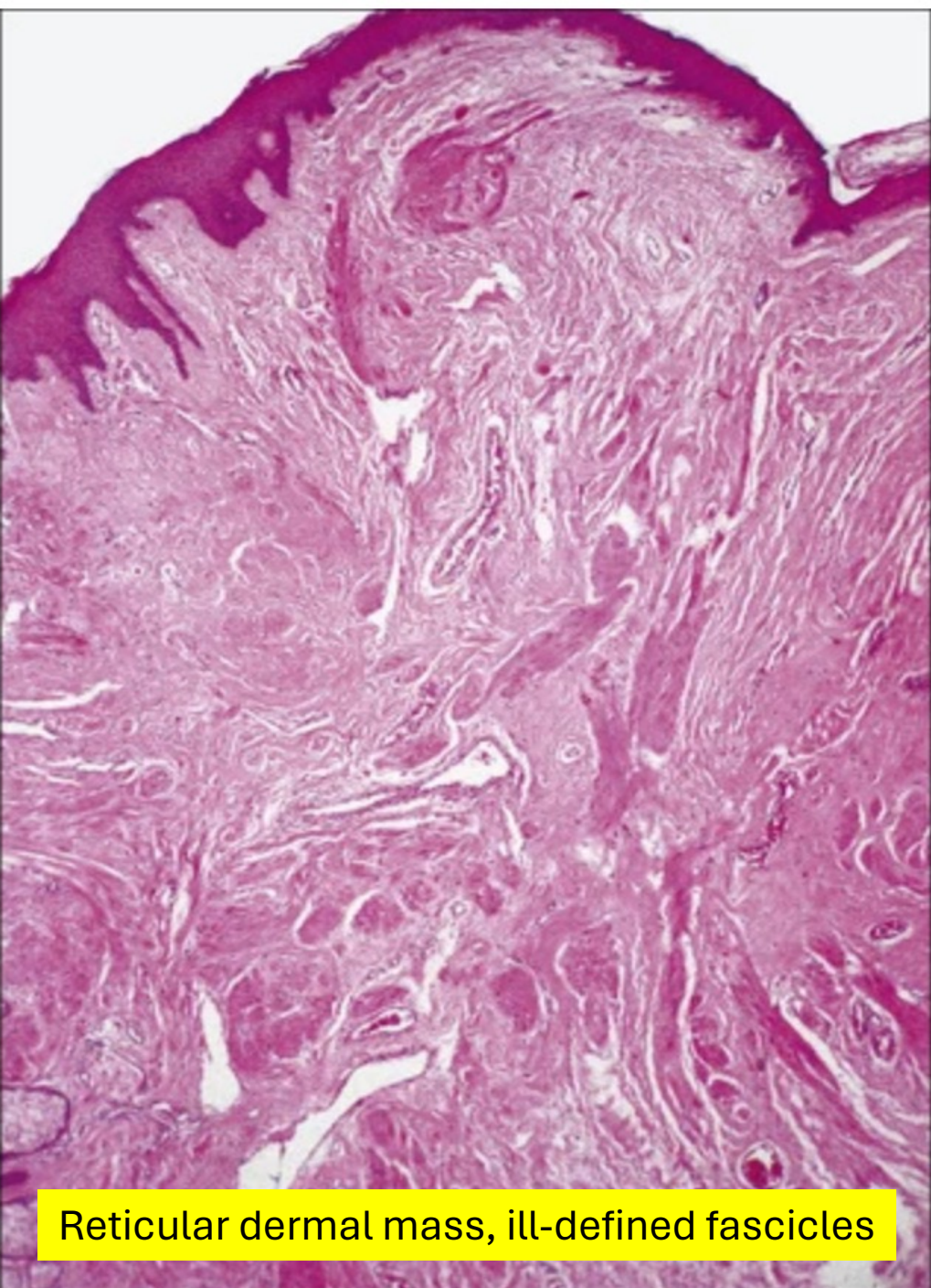
Bundles of eosinophilic cells with cigar-shaped nuclei

Reticular dermal mass, ill-defined fascicles

PILAR LEIOMYOMA

IHC: SMA+, desmin+, and h-caldesmon+, S100A6-

- Young adults, limbs or trunk
- Can present as multiple papules, painful/tender (cold or compressed)
- Hereditary leiomyomatosis and renal cell cancer (HLRCC), *FH* (1q42.3~q43):
 - Cutaneous and uterine leiomyomas, renal cell carcinoma
- Fumarate hydratase deficiency (tricarboxylic acid cycle):
 - TS gene and DNA damage response
- DDX: DF, dermatomyofibroma, cellular neurofibroma, leiomyosarcoma (atypical intradermal smooth muscle neoplasm: less mitoses and nuclear pleomorphism)

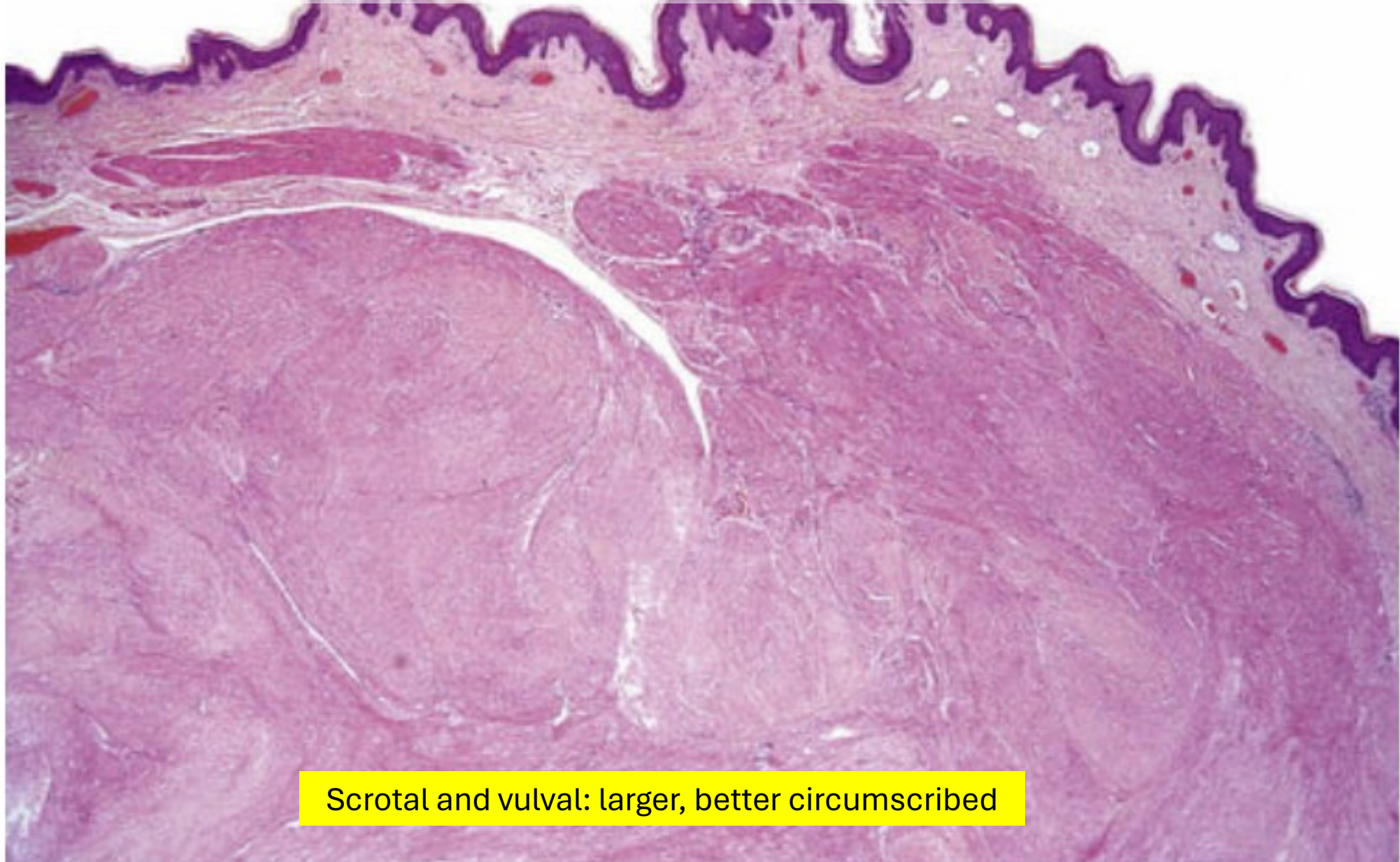


Reticular dermal mass, ill-defined fascicles

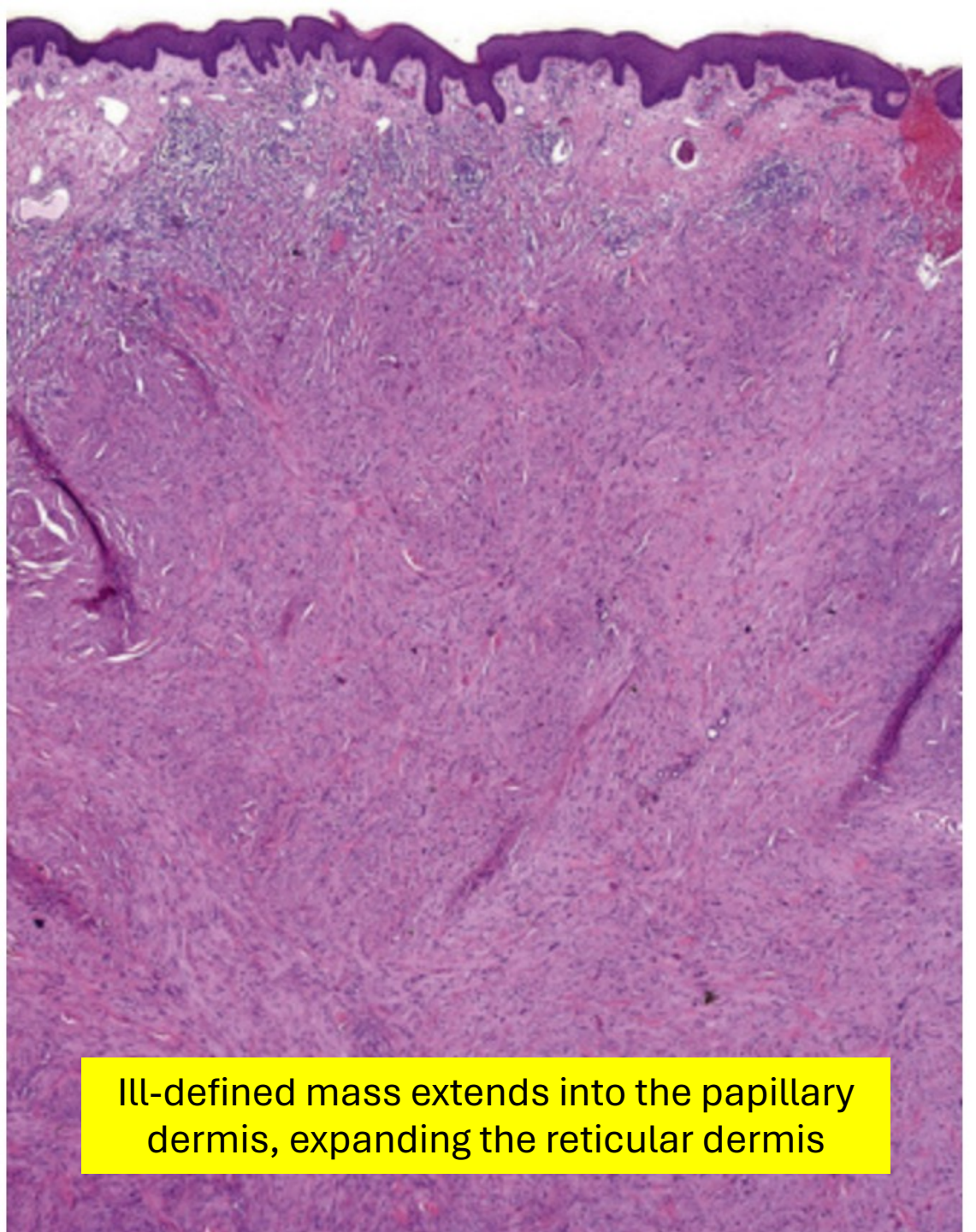
GENITAL LEIOMYOMA

- Originates from the smooth muscle:
 - scrotum (dartos muscle)
 - vulva (labia majora) or
 - nipple
- Middle-aged adults
- Multiple leiomyomas of the vulva (Alport syndrome)

SCROTAL LEIOMYOMA



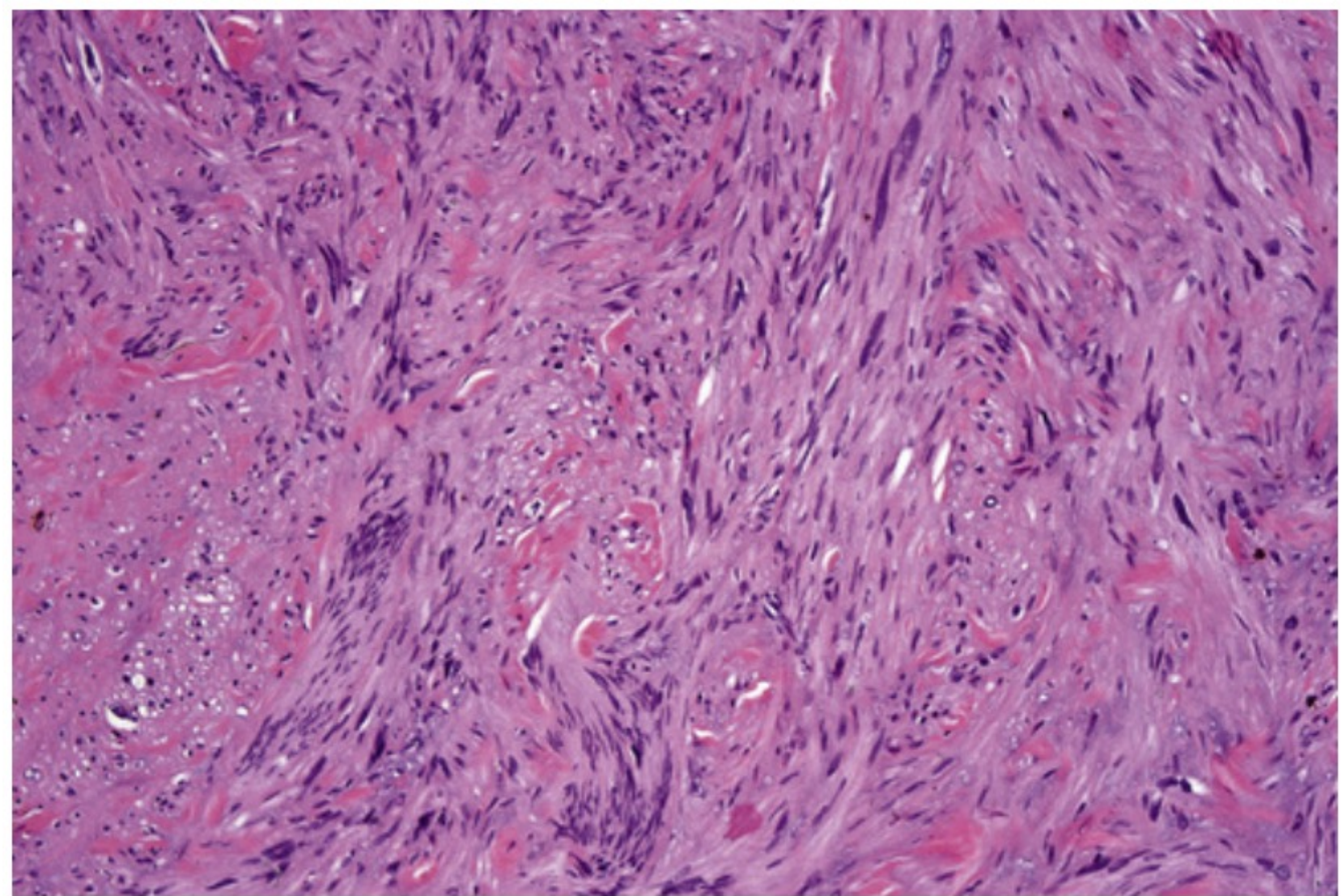
Scrotal and vulval: larger, better circumscribed



Ill-defined mass extends into the papillary dermis, expanding the reticular dermis

LEIOMYOSARCOMA

- Deep: abdomen and retroperitoneum
- Superficial:
 - Cutaneous: leiomyosarcoma of the nipple
 - Arrector pili origin
 - Middle-aged males
 - Trunk and limbs
 - Local recurrence common
 - Margin status critical (> 1 cm)
 - Mohs with follow-up
 - Subcutaneous: scrotal and vulval variants related to deep
 - Vein wall origin
 - Older adult males, thighs
 - Larger than cutaneous lesions
 - Local recurrence common
 - 50% metastasize (mortality 30-50%)
- Metastasis rare (to skin)



Eosinophilic spindled cells with hyperchromatic and pleomorphic nuclei

Mitotic activity

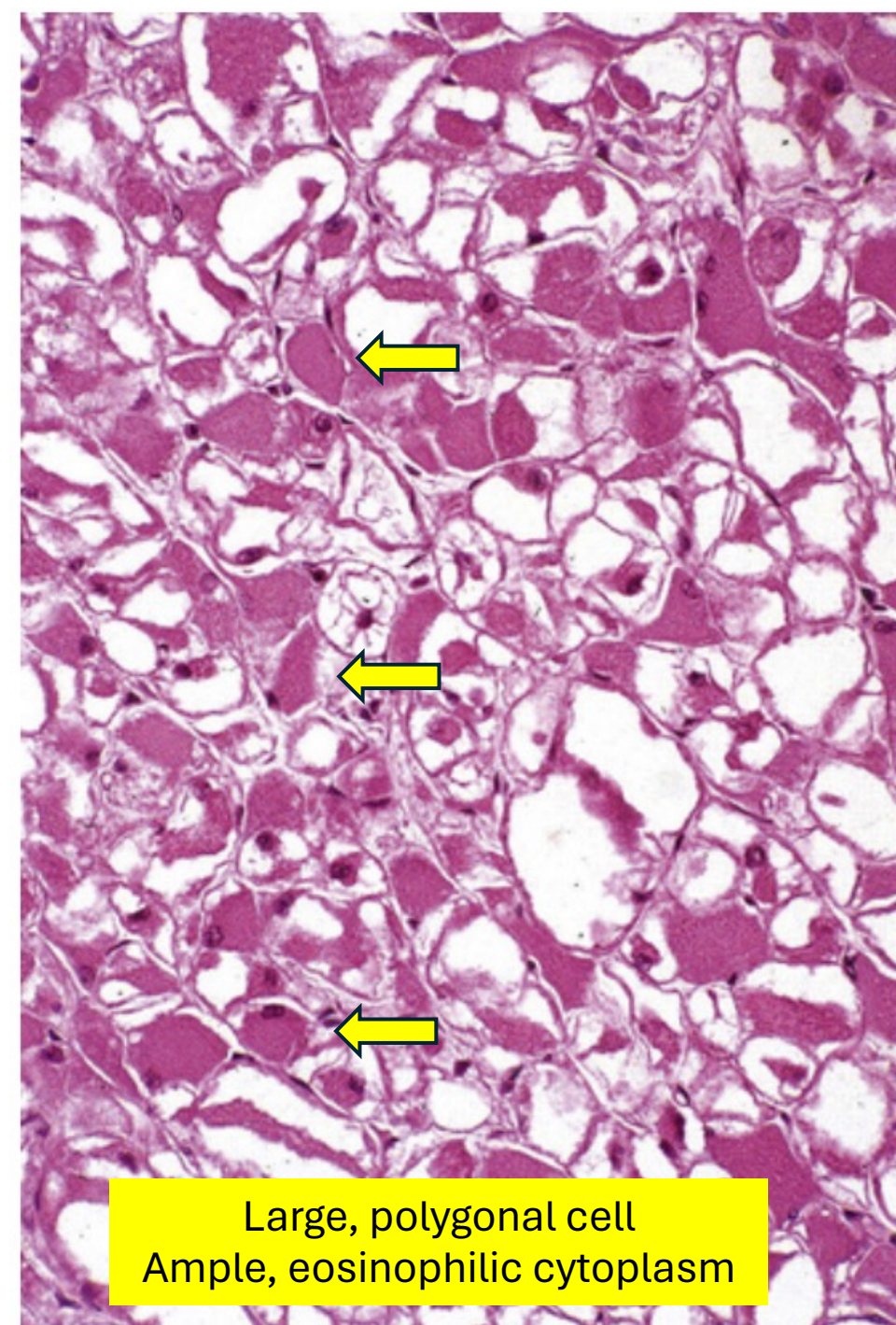
Shares eosinophilia with leiomyoma

LEIOMYOSARCOMA

- IHC: SMA+, desmin+, and h-caldesmon+
 - PTEN- (loss)
 - P53+, S100A6+
 - ki-67 increased proliferative index
- Epithelioid variant exists
- Malignant features: necrosis and hemorrhage
- DDX: spindle cell melanoma (S100+), metastatic leiomyosarcoma (needs CPC)

RHABDOMYOMA (EXTRACARDIAC)

- Rare, deep-seated mass
- Genital type:
 - Middle-aged women
 - Vagina, cervix > vulva
 - Males: paratesticular soft tissue
- Adult type:
 - Older adult males
 - Head and neck, oral cavity
- Fetal type:
 - Male infants
 - Face, neck
- IHC: muscle-specific actin+, myoglobin+, desmin+
- DDX: rhabdomyosarcoma

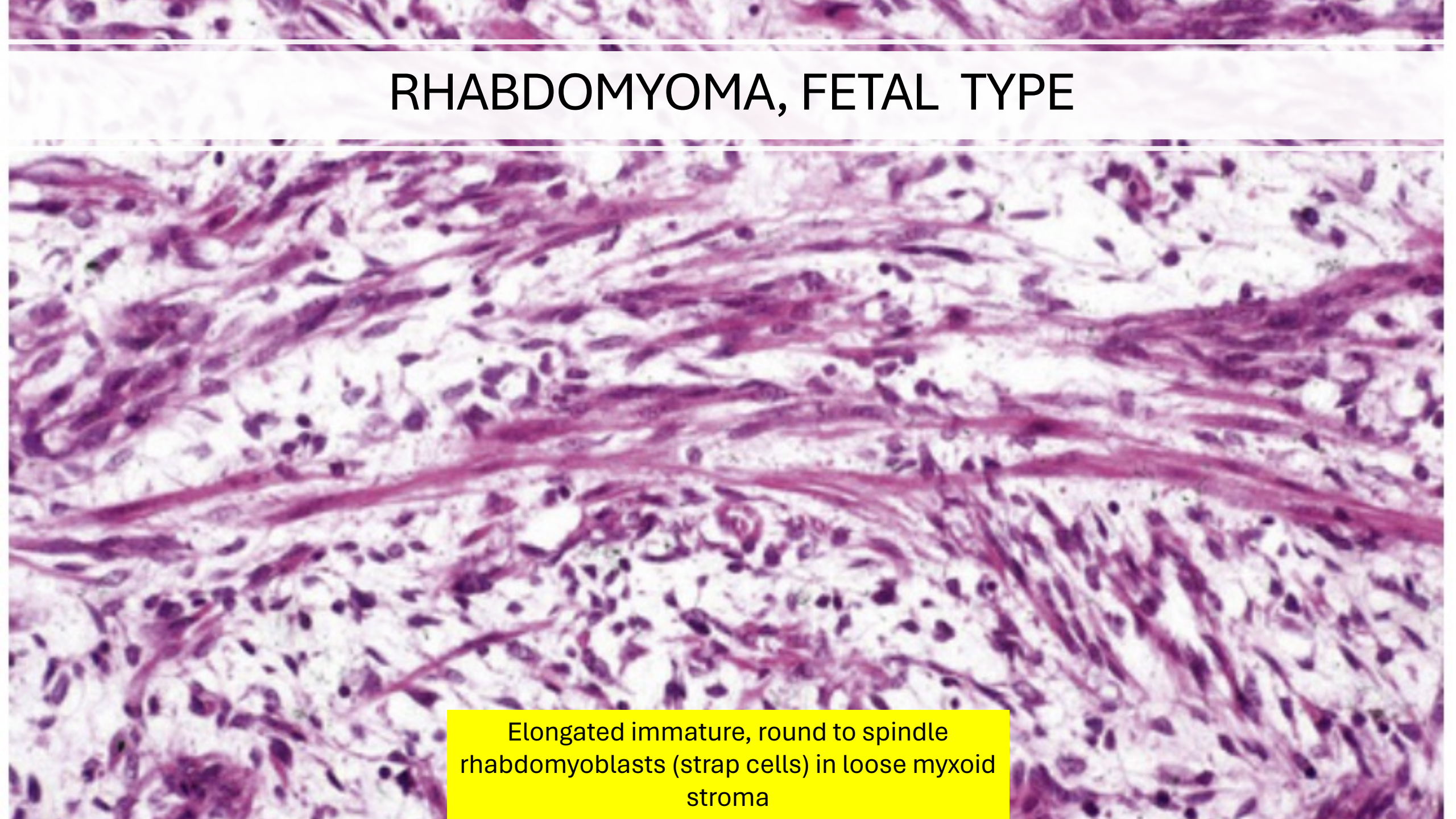


RHABDOMYOMA, ADULT TYPE



'jack straw' cross-striations, intracytoplasmic inclusions

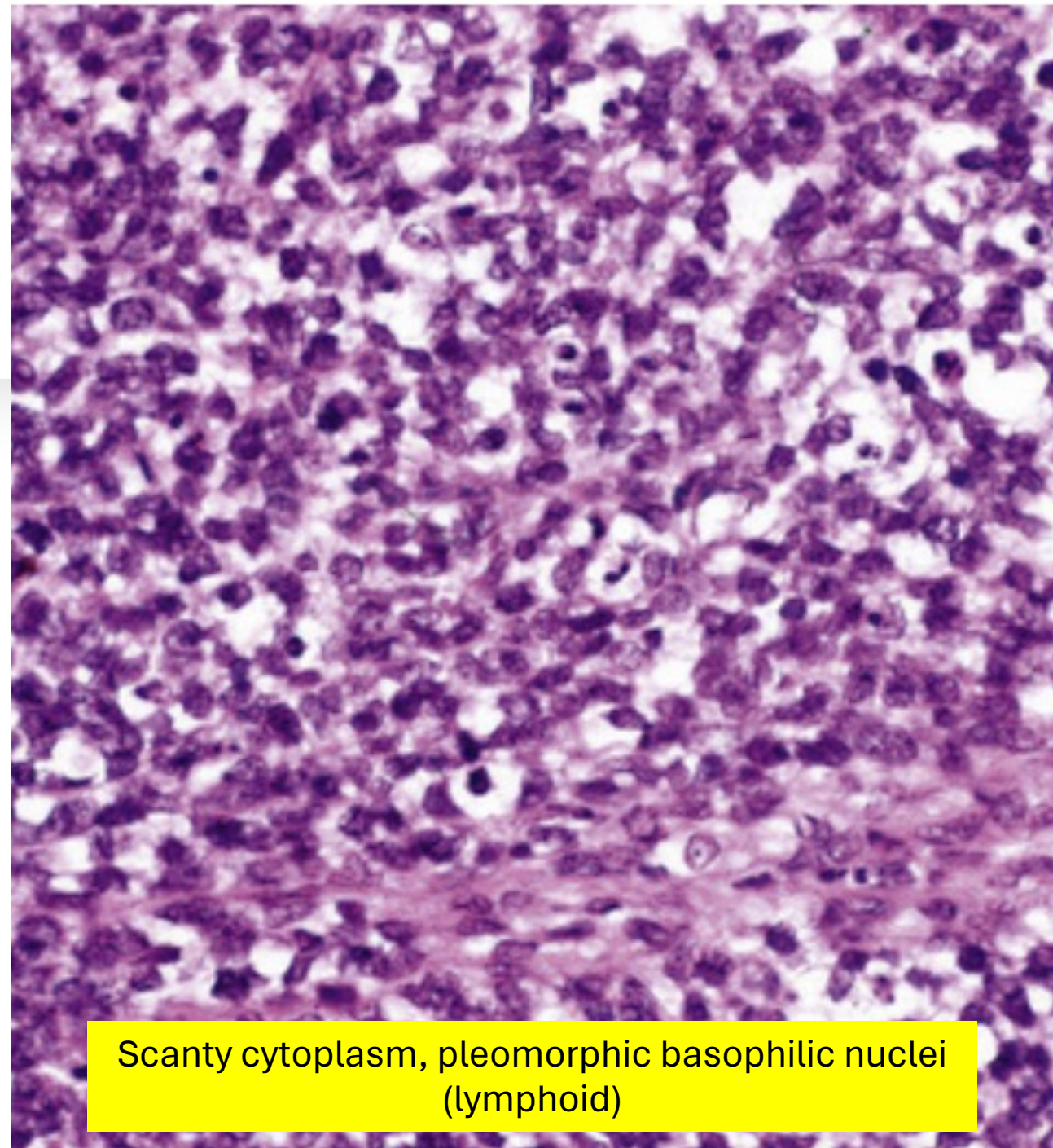
RHABDOMYOMA, FETAL TYPE



Elongated immature, round to spindle rhabdomyoblasts (strap cells) in loose myxoid stroma

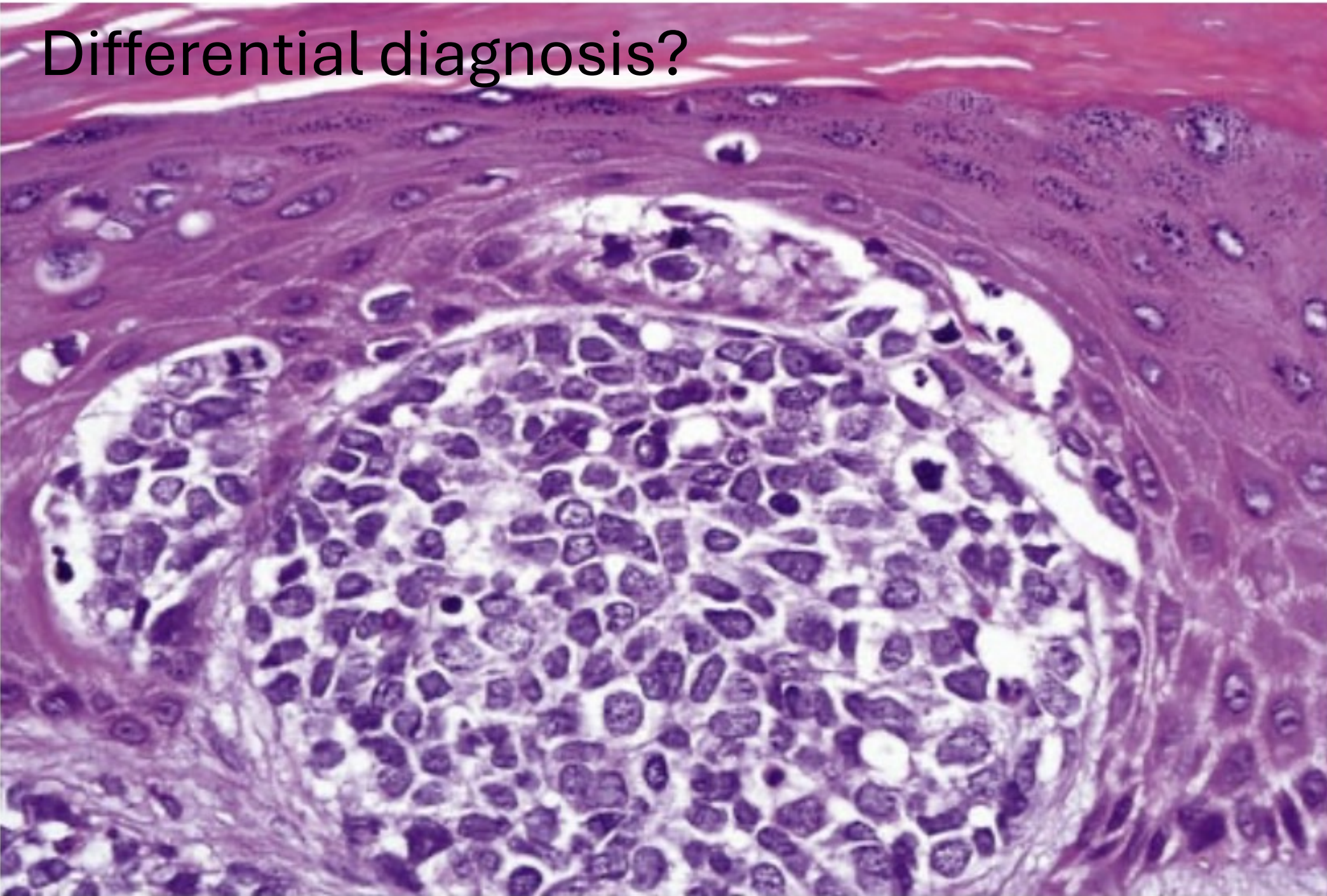
RHABDOMYOSARCOMA

- Very rare
- Bimodal age (mean): 10 and 74 years
- 4 histologic types:
 - Alveolar and embryonal (peds)
 - Pleomorphic and spindle cell/sclerosing (adults)
- Variable genetic alterations
- IHC: desmin+, MSA+, myogenin+, MyoD1
- DDX (small round cell tumor): neuroblastoma, primitive neuroectodermal tumor, MCC, lymphoma, melanoma



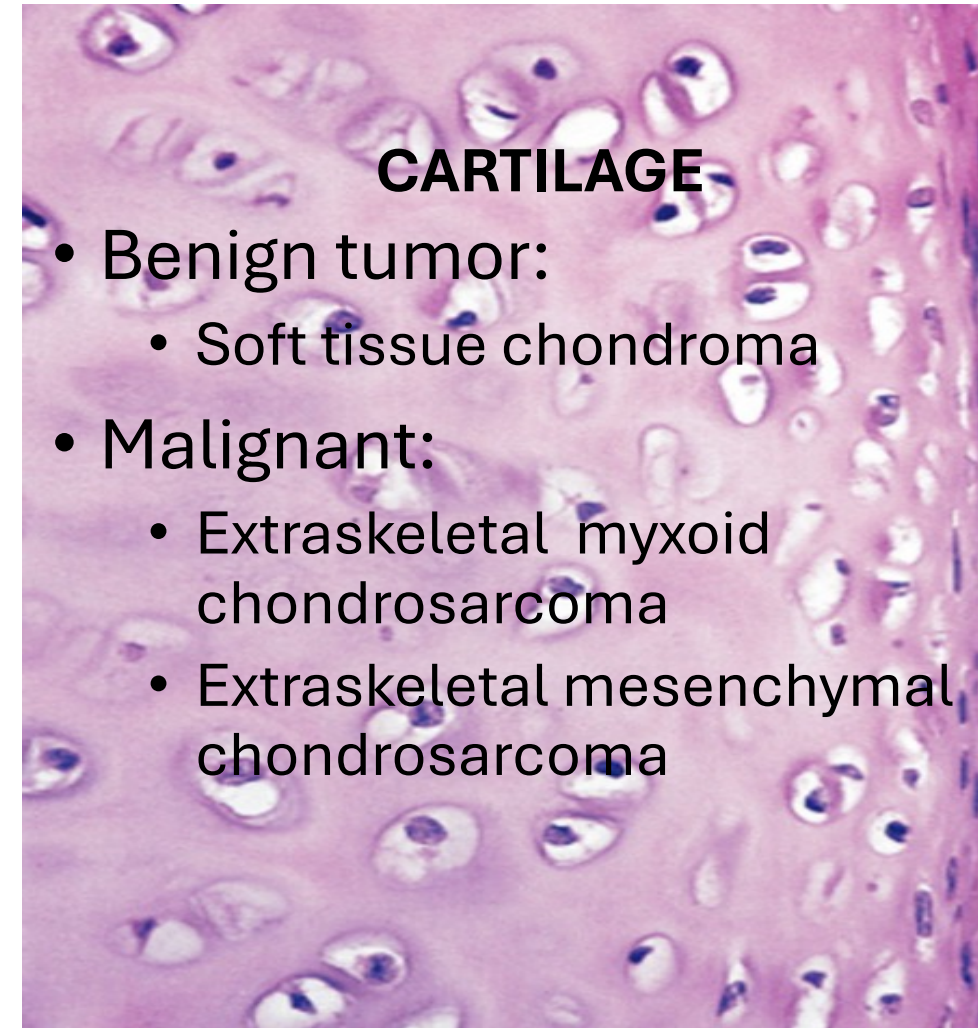
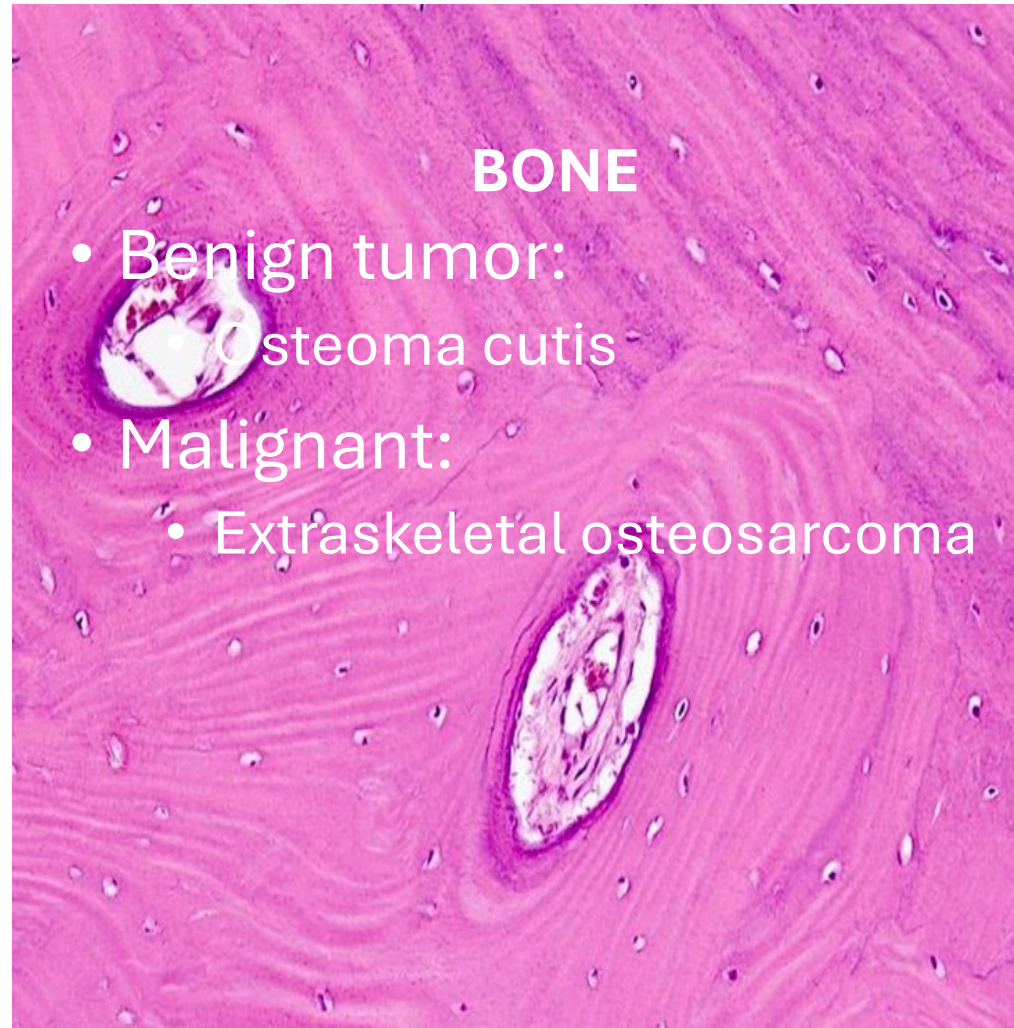
Scanty cytoplasm, pleomorphic basophilic nuclei (lymphoid)

Differential diagnosis?



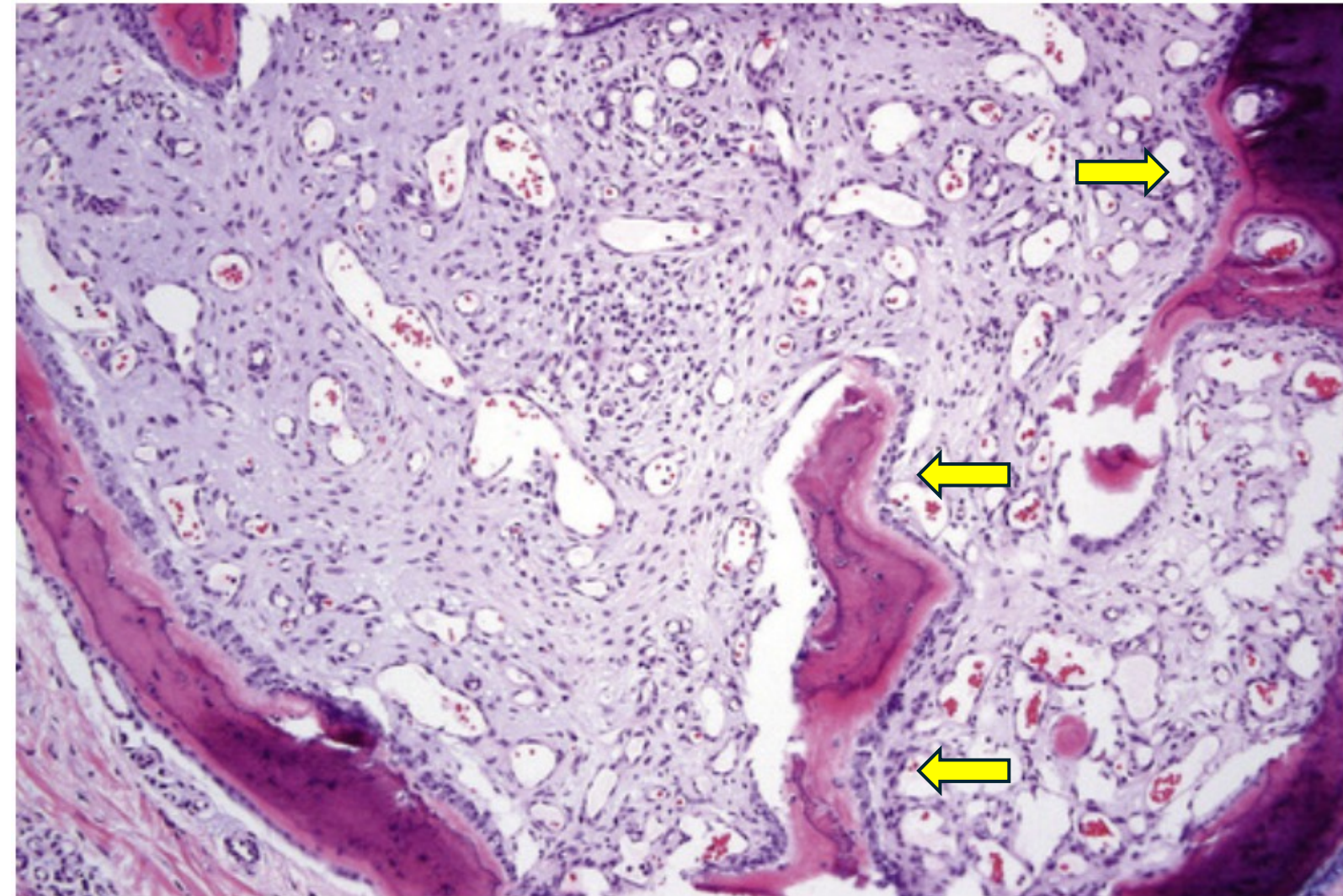
Alveolar rhabdomyosarcoma

TUMORS OF BONE AND CARTILAGE



TUMORS OF BONE AND CARTILAGE

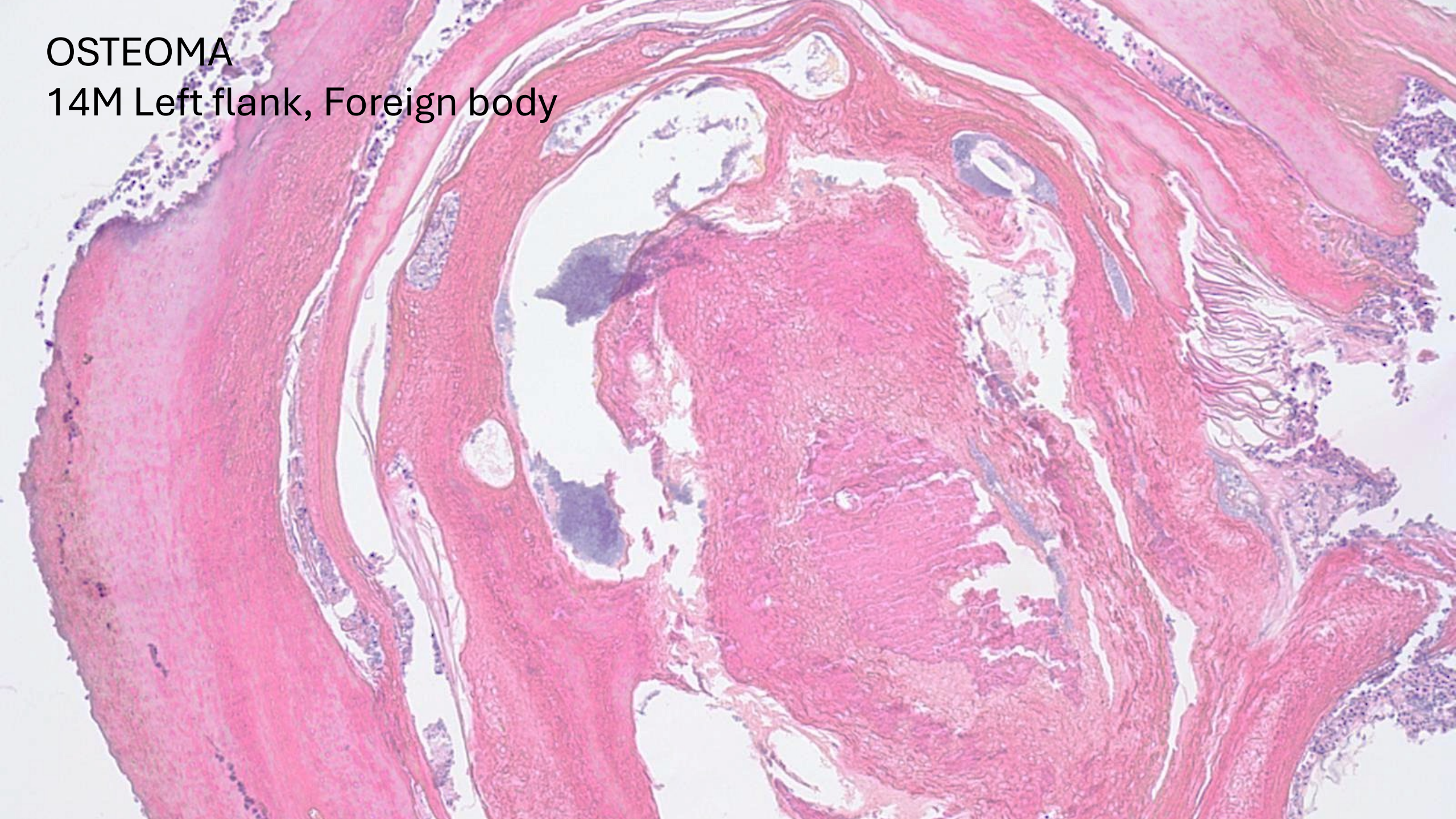
- Majority of ossification in skin:
 - Secondary degenerative, e.g., OC
 - Metaplastic process
 - Melanocytic nevi
 - Pilomatrixoma
- Primary lesions are very rare
- Osteoma cutis-
 - Dystrophic ossification in acne and folliculitis
 - Syndromes: e.g., Albright hereditary osteodystrophy
- DDX: benign cartilaginous exostosis (osteochondroma)



Bone formation, osteoid area rimmed by osteoblasts (ant trails), scarred medullary cavity

OSTEOMA

14M Left flank, Foreign body

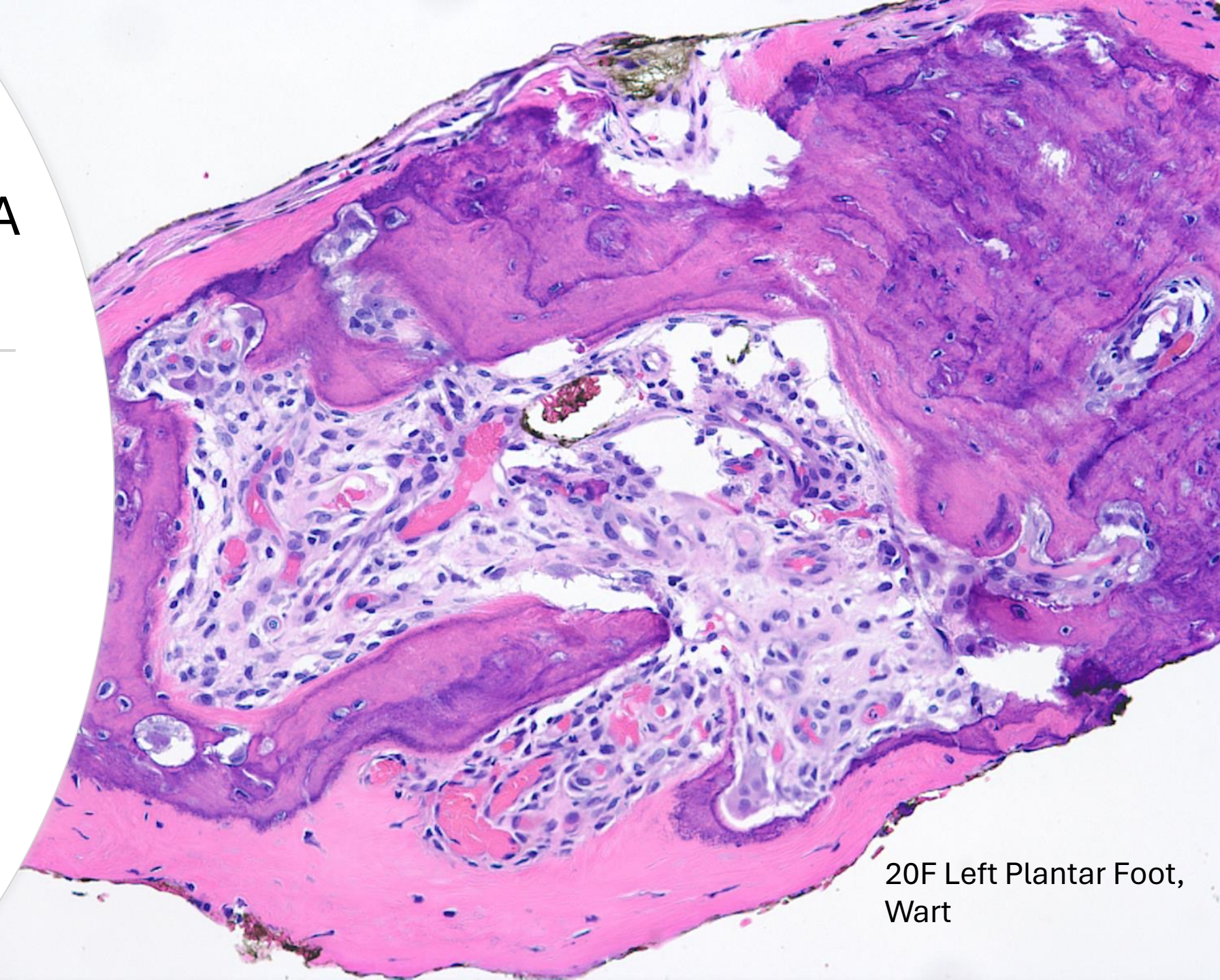




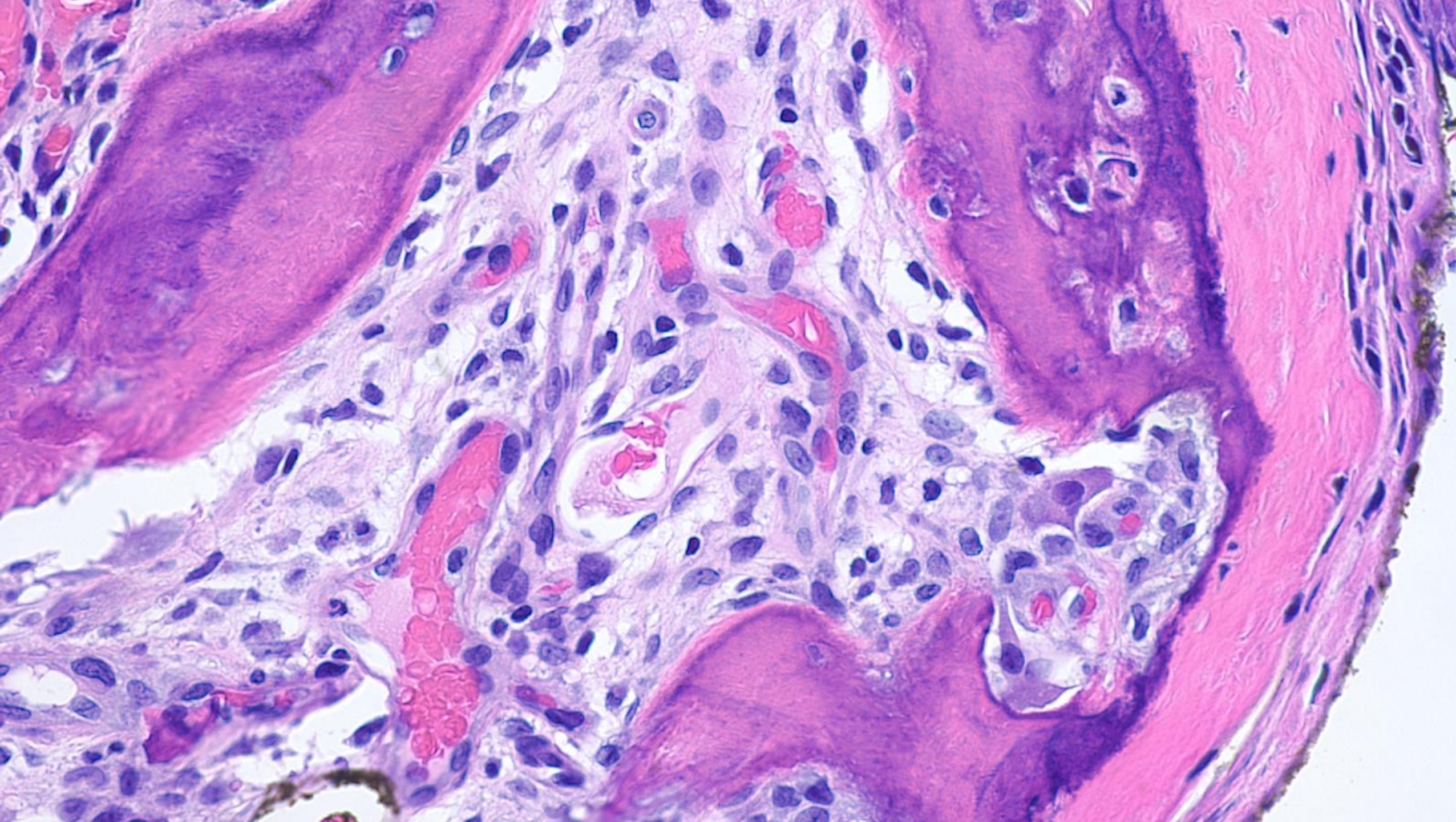
Medullary cavity, containing bone marrow cells

OSTEOCHONDROMA (EXOSTOSIS)

- Solitary, under the nail, hard and painful tumor
- Mature cartilage overlying a layer of lamellar bone
- Arises from the underlying phalanx

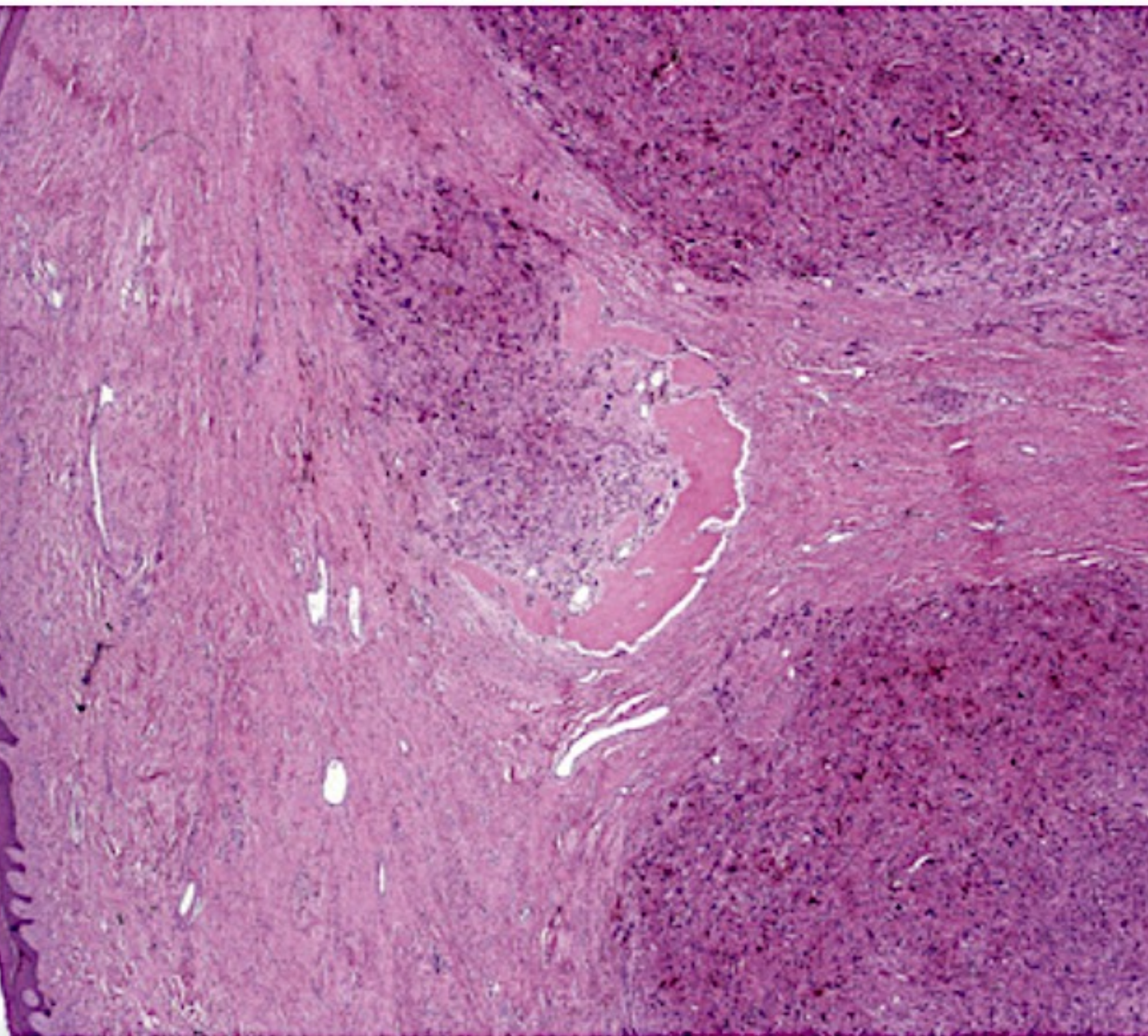


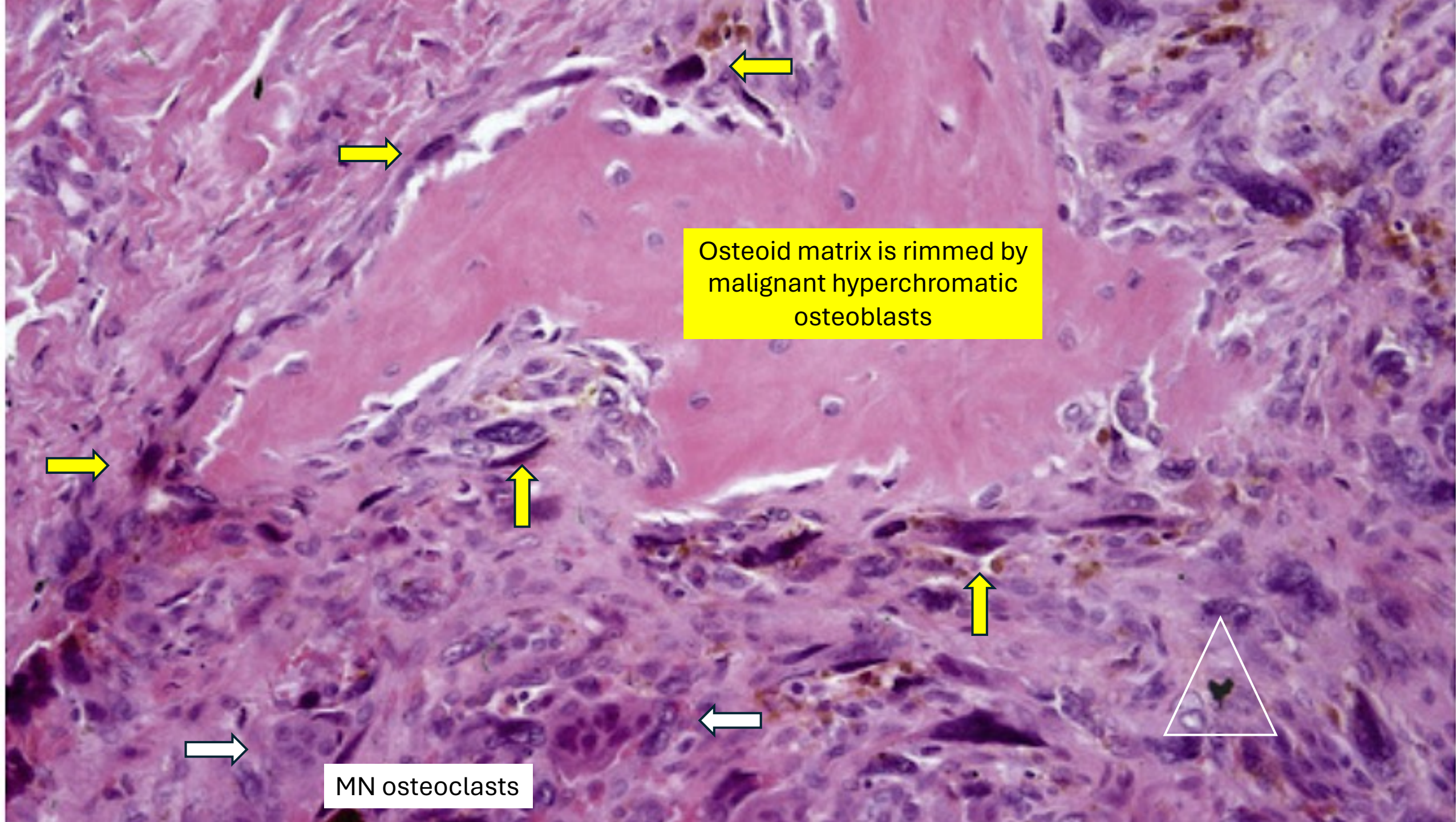
20F Left Plantar Foot,
Wart



EXTRASKELETAL OSTEOSARCOMA

- Rare, in older adults
- Unusual in children
- Deep soft tissue, subcutis or dermis of the limbs (legs)
- 10% associated with XRT
- Rapid local recurrence, widespread metastasis with 75% mortality
- 12q amplification better prognosis
- Loss of CDKN2A biallelic simultaneous losses of RB1 and TP53
- IHC: Osteocalcin+, SATB2+, ERG (cartilaginous area)





MN osteoclasts

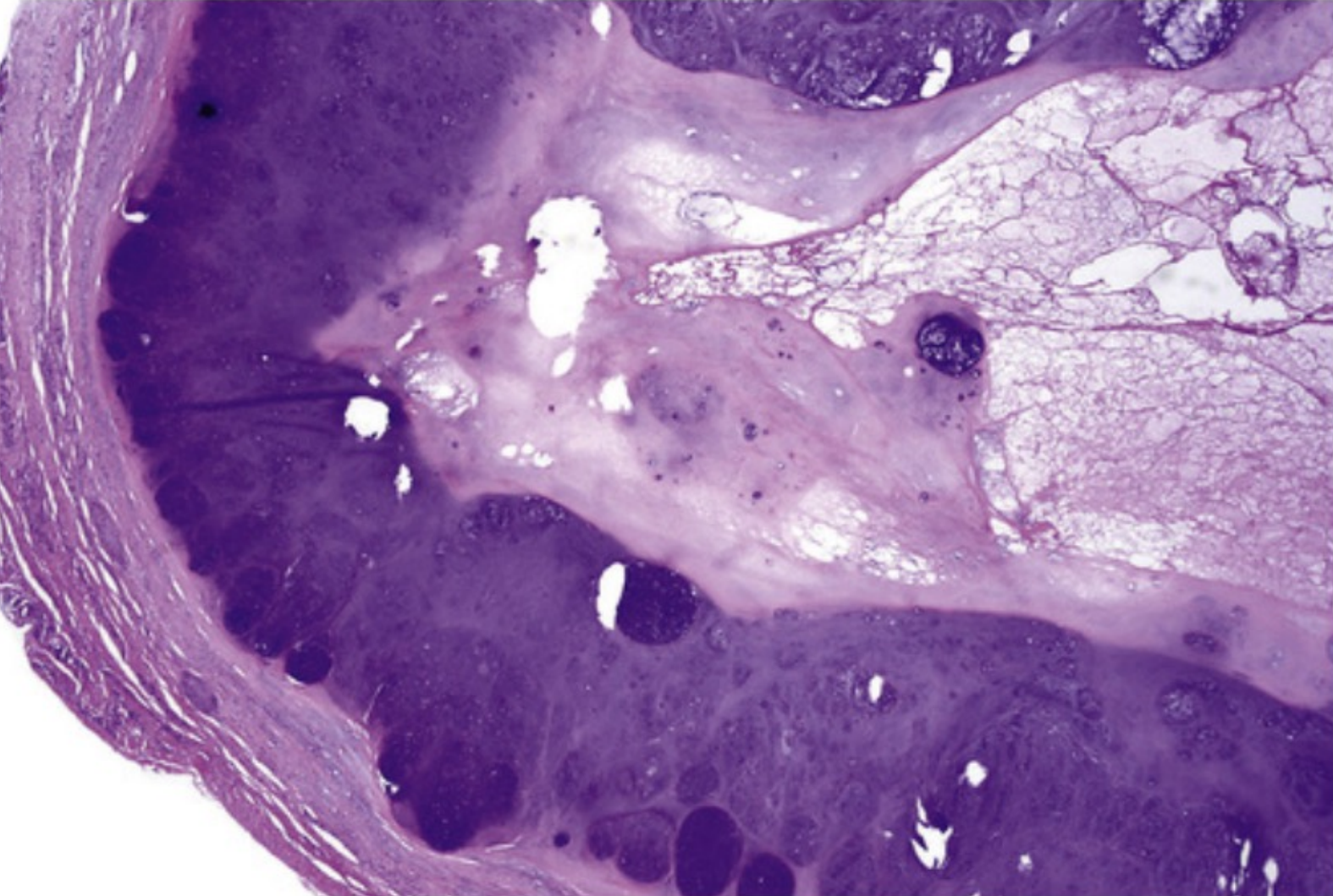


Osteoid matrix is rimmed by malignant hyperchromatic osteoblasts



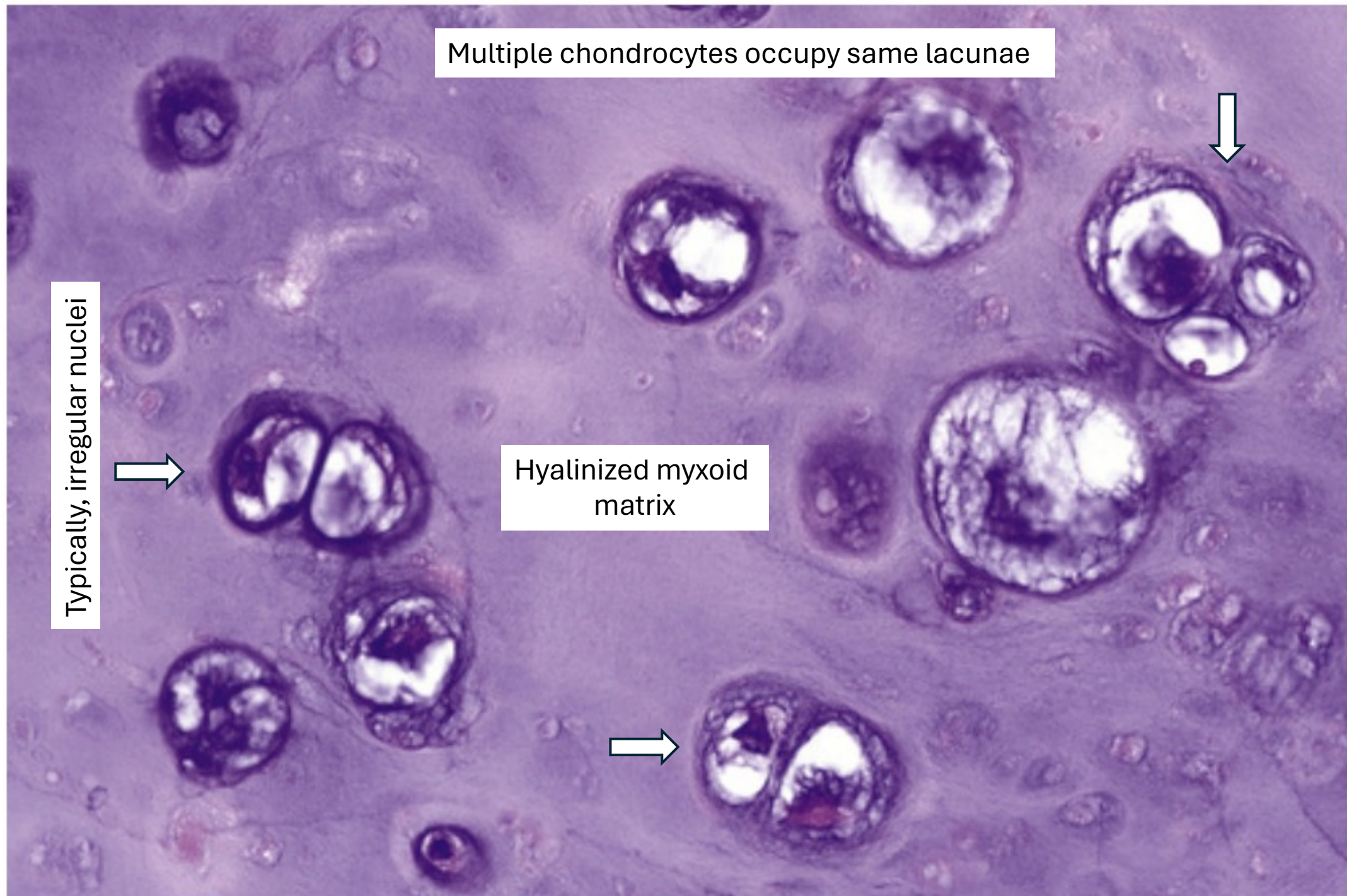
SOFT TISSUE CHONDROMA

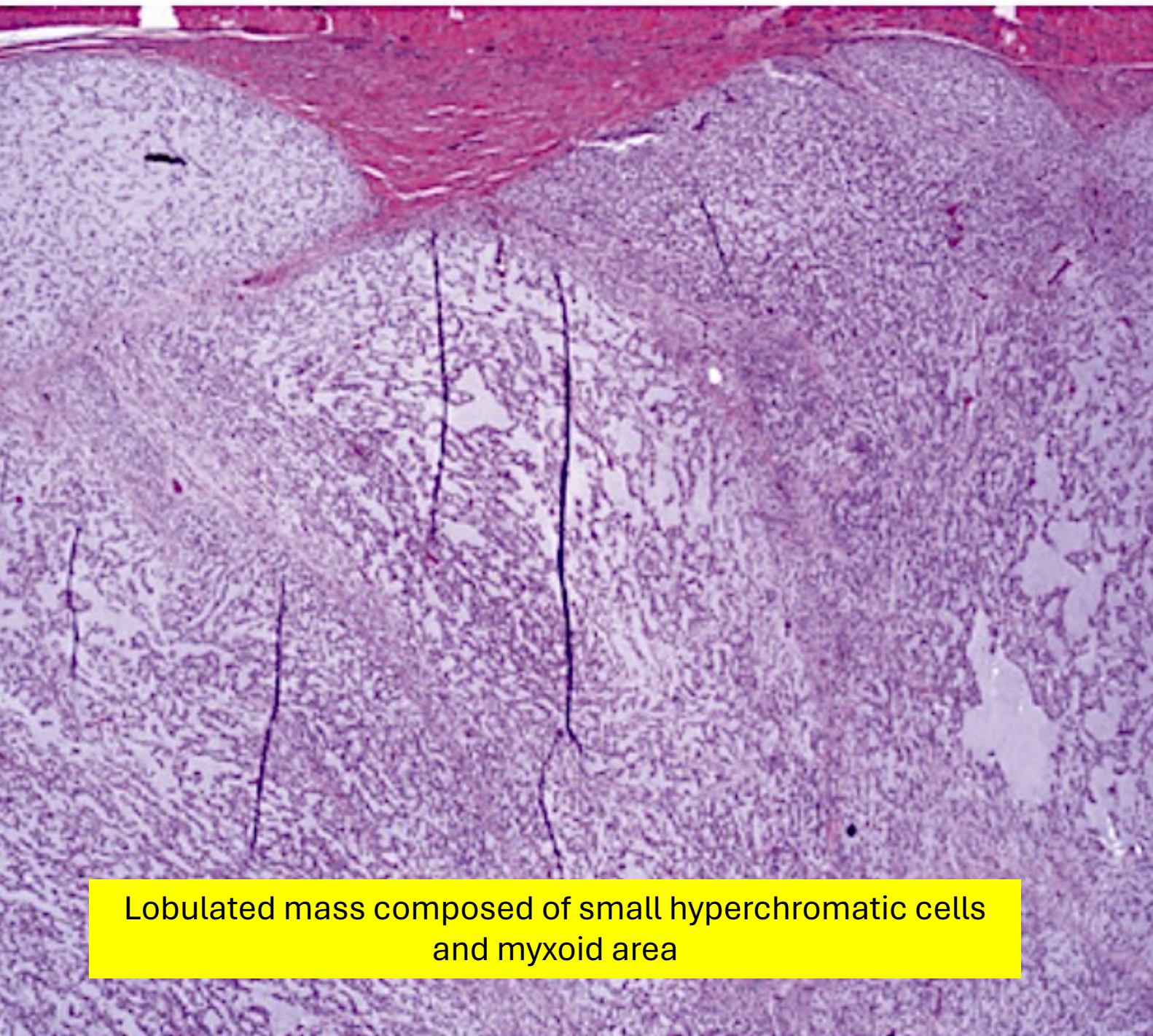
- Uncommon, middle-aged males on hands or feet
- 10% recurrence, no malignant change
- 12q 13-15 rearrangements, expression of *HMGA2*
- Monosomy 5, trisomy 6
- Histology: degenerative changes: myxoid, hemorrhage, calcification or ossification
 - Reactive osteoclastic giant cells
- IHC: ERG (cartilaginous area)
- DDX: mixed tumor of skin



Intradermal, well-circumscribed, lobulated mass of mature hyaline cartilage

SOFT TISSUE CHONDROMA





Lobulated mass composed of small hyperchromatic cells and myxoid area

EXTRASKELETAL MYXOID CHONDROSARCOMA

- Uncommon, adult males (legs)
- Deep origin, 20% subcutaneous
- Recurs and metastasizes
 - 5-, 10- & 15-year survival: 82%, 65% and 58%
- t(9;22) (q22;q12) fusing *NR4A3* and *EWSR1*
- IHC: INI1 loss, S100+, SOX10+, rare EMA+, Keratin+
- DDX: myxoid liposarcoma and malignant mixed tumors

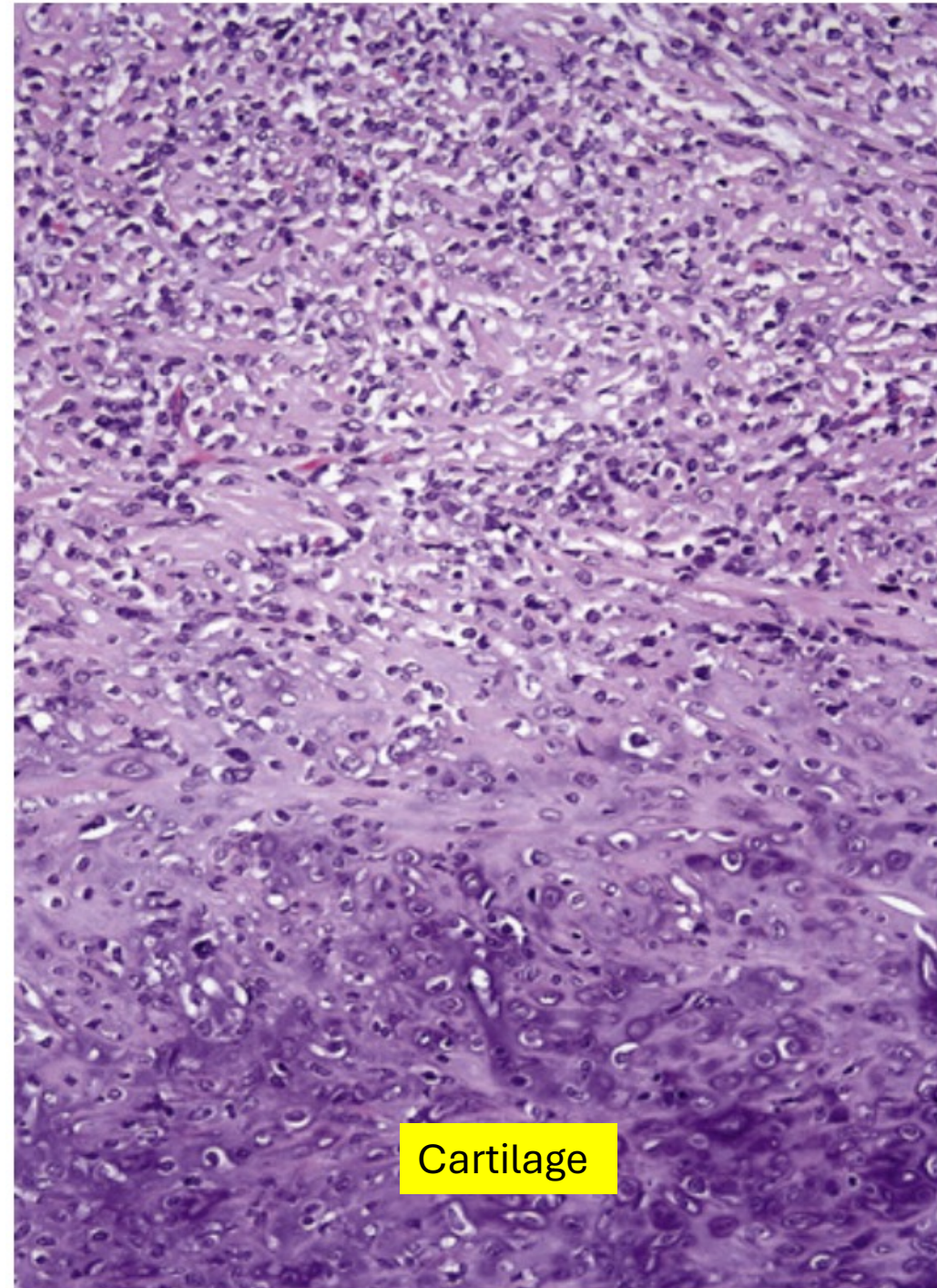
EXTRASKELETAL MYXOID CHONDROSARCOMA

A light micrograph showing a tissue sample with a mix of cellular and fibrous components. In the center, there is a cluster of cells with dark, hyperchromatic nuclei and some clear, vacuolated areas. The surrounding tissue appears more fibrous and less cellular.

Plump spindle cells with pleomorphic
and hyperchromatic nuclei

EXTRASKELETAL MESENCHYMAL CHONDROSARCOMA

- Very rare, deep tumor in young adults, children (females)
- Head and neck, upper trunk > limbs
- Prognosis worse than the myxoid variant
- Inv(8) (q13q210), *HEY1-NCOA2* fusion
- IHC: CD99+, BCL-2+, S100+, ERG+, SOX-9+
- Biphasic Histology: hypercellular tumor with undifferentiated round or spindled mesenchymal cells AND mature cartilage



References

- McKee's *Pathology of the Skin* Eduardo Calonje

McKEE'S
Pathology of the Skin
WITH CLINICAL CORRELATIONS



Eduardo Calonje

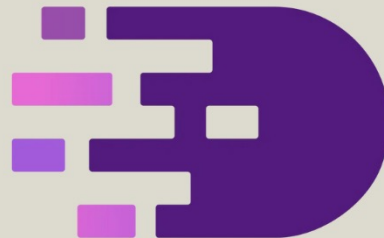
Thomas Brenn | Alexander J. Lazar | Steven D. Billings



Digital Skin Pathology

<https://digitalskinpathology.com/>

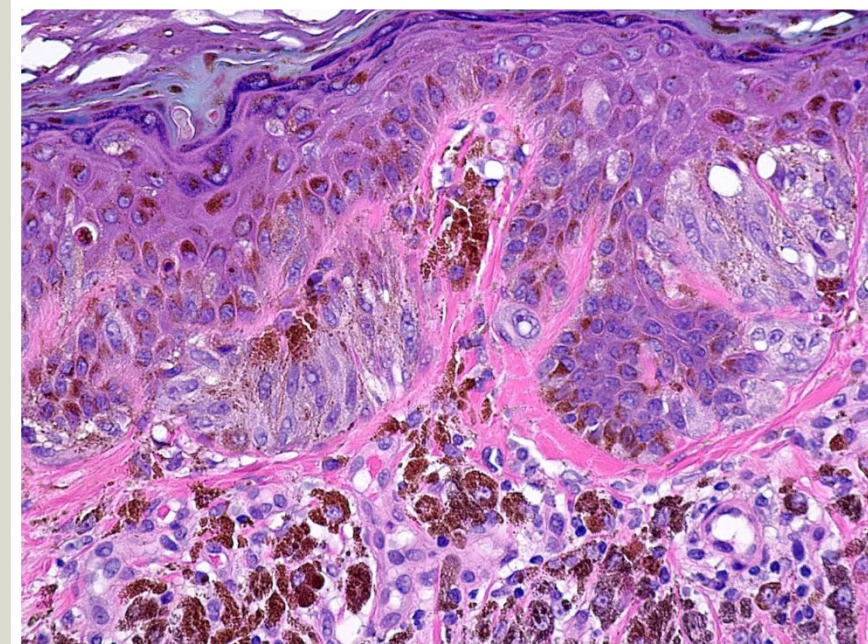
- Current lecture
- Examples of cases



DIGITAL SKIN PATHOLOGY (DiSK)

Learn Histologic Diagnosis Case-By-Case

**DERMATOPATHOLOGY: LEARN HOW TO
DIAGNOSE SKIN DISEASES DERM PATH
DIAGNOSTICS**



Understand your patient's dermatopathology diagnostic report to provide better clinical care (how to diagnose skin diseases), derm path diagnostics