Mycosis Fungoides
Definition

- Mature T-cell lymphoma
- Presents in skin with patches/plaques
- Characterized by epidermal and dermal infiltration of small to medium-sized T-cells with cerebriform nuclei
Epidemiology

- Most common primary T-cell lymphoma of skin
- Incidence 0.29/100,000/year
- 0.5% of NHL
- Adults/elderly
- M:F = 2:1
Sites of involvement

- **Early stages**
  - Limited to skin

- **Advanced stages**
  - Skin
  - Extracutaneous dissemination
    - LN
    - Liver
    - Spleen
    - Lungs
    - PB
    - BM (rare)
Clinical features

- Long natural history

- Non-specific scaly eruptions years before diagnostic histology develops
Clinical features

- Initial diagnostic lesions
  - Limited patches and/or plaques
    - Frequently on trunk
    - May persist for years

- Later diagnostic lesions
  - More generalized plaques
  - Tumors
Mycosis Fungoides

Early stage

Plaque stage

Tumor stage
Mycosis Fungoides
Tumor Stage
Clinical features

- Rare patients may develop generalized disease with erythroderma
  - May overlap with Sézary syndrome

- Extracutaneous dissemination
  - Late event
  - Predominantly in patients with extensive/advanced cutaneous disease
Clinical features

- D’emblée lesions
  - Skin tumors without a preceding patch/plaque stage
  - Rare
  - “not entirely well defined”
  - May represent other subtypes of T-cell lymphoma with preferential cutaneous infiltration
Etiology

- Unknown pathogenesis

- HTLV-1 (or similar virus) implicated in one series (Pancake et al. 1995)
  - 50 patients
  - Truncated proviral sequences similar to tax and/or pol detected by PCR in 30-90%
  - Majority of patients have antibodies to tax, but not to the structural proteins of HTLV-1
  - Whether the virus is the cause or secondary event is unknown
  - Not identified in a European series
Morphology

- Epidermotropism
  - Small to medium-sized cells with irregular (cerebriform) nuclei
  - Larger cells with similar nuclei (minority)
  - Involvement with single cell exocytosis is most common form of epidermotropism
Morphology

- Pautrier microabscesses
  - Only seen in a proportion of cases
  - Highly characteristic
Mycosis Fungoides

- Epidermis with Pautrier microabscess
- Atypical lymphoid cells
Mycosis Fungoides
Pautrier Microabscesses
Morphology

- Dermal infiltrates
  - Patchy
  - Band-like
  - Diffuse
  - Often associated with inflammatory infiltrate of small lymphocytes and eosinophils
Morphology

- **Patch**
  - Sparse infiltrate of lymphocytes spread along the papillary dermis
  - Only slight cytological atypia
Mycosis Fungoides
Morphology

- Plaque
  - More dense infiltrate of atypical lymphocytes that can extend around the adnexae
  - Atypical lymphocytes are more common
    - 10-30 μm in diameter
    - Prominent nuclear convolutions (cerebriform)
Morphology

- Tumor
  - Involvement of entire dermis +/- subcutis
  - Infiltrate of larger atypical lymphocytes
Morphology

- LN involvement
  - Dermatopathic lymphadenopathy
    - Paracortical expansion due to the presence of large number of histiocytes and interdigitating cells
Morphology

- LN involvement, category II
Morphology

- LN involvement
  - Clonal T-cells present in
    - Majority of category II and III lesions
    - Occasionally in category I lesions
  - Presence of clonal T-cells may be associated with unfavorable outcome
Immunophenotype

- CD2/3/4/5 and TCRβ positive
- HECA antigen (associated with lymphocyte homing to skin) positive in most cases
- CD7/8 negative
- Cytotoxic granule associated proteins negative in early patch/plaque lesions
Mycosis Fungoides

CD3
Immunophenotype

CD3

CD4 negative
(most cases positive)

CD8
Genetics

- TCR clonally rearranged
- Inactivation of CDK2A/p16 and PTEN identified in two studies
- Complex karyotypes present in many patients (particularly in advanced stages)
- No specific chromosomal abnormality identified
Prognosis

Most important prognostic factor is clinical stage

Limited disease

- Excellent prognosis
- Survival similar to general population

Advanced stages: poor prognosis, especially with

- Skin tumors
- Extracutaneous dissemination
Prognosis

- Other adverse prognostic indicators
  - Age > 60 years
  - Elevated LDH
  - Transformation to a large T-cell lymphoma
MF Variants

- Pagetoid reticulosis
  - Infiltrate is strictly epidermal
MF Variants

- Pagetoid reticulosis

- Distinguish between
  - Woringer-Kolopp disease (localized skin lesions)
  - Ketron-Goodman disease (multiple skin lesions)

[It is generally recommended that the designation be restricted to the localized variants which have an excellent prognosis]
MF Variants

- **Pagetoid reticulosis**
  - Cerebriform neoplastic T-cells
  - Often CD30 positive
  - CD4+/CD8- or CD4-/CD8+ or CD4-/CD8-
  - TCR rearrangement in some cases
MF Variants

- MF-associated follicular mucinosis
  - Rare
  - Follicular (rather than epidermal) infiltrates of cerebriform T-cells
  - Mucinous degeneration of hair follicles
MF Variants

- MF-associated follicular mucinosis
  - Head and neck
  - TCR clonal rearrangement in most cases
  - Indolent (but slightly less favorable prognosis than MF)
Mycosis Fungoides-associated Follicular Mucinosis
Mycosis Fungoides-Associated Follicular Mucinosis
Mycosis Fungoides
Granulomatous Slack Skin Disease

- May be seen with other types of lymphoma (e.g., Hodgkin) and may be separate entity
- Slowly developing folds of atrophic skin
- Preferentially involves axillae or groin
- Granulomatous infiltrate with atypical T lymphocytes, macrophages, multinucleated giant cells
- Elastophagocytosis
Granulomatous Slack Skin Disease
Sézary Syndrome
Definition

- Generalized mature T-cell lymphoma

- Characterized by
  - Erythroderma
  - Lymphadenopathy
  - Neoplastic T-lymphocytes in PB (cerebriform)

- MF variant, but behavior is much more aggressive
Epidemiology

- Rare
- Exclusively in adults
Sites of involvement

- Generalized disease with involvement of
  - Skin
  - LN
  - PB
  - Visceral organs, in the terminal stages

- BM often spared
Clinical features

- Patients present with
  - Erythroderma
  - Generalized lymphadenopathy
  - Pruritis
  - Alopecia
  - Palmar or plantar hyperkeratosis
  - Onychodystrophy
Sezary Syndrome

Diffuse erythroderma
Sezary Syndrome
Diffuse Erythroderma
Etiology

- Unknown pathogenesis
- Association with HTLV-1 is controversial
Morphology

- Skin lesions are similar to MF with dermal and epidermal infiltrates of cerebriform T-lymphocytes

- LN
  - Effaced architecture
  - Paracortical or diffuse infiltrates
  - With or without changes of dermatopathic lymphadenopathy
Morphology

- Neoplastic cells in PB
  - Markedly convoluted nuclei
  - Predominantly small (Lutzner cells), or large (classical Sézary cells), or mixture of both
Sezary syndrome

Cells with convoluted nuclei

Cebebriform nuclei (EM)
Sezary Syndrome

Sezary cells in peripheral blood
Sezary Syndrome
Morphology

- Neoplastic cells in PB
  - No consensus on degree of lymphocytosis
  - Most studies require ≥1000 Sézary cells per mm$^3$
Morphology

- BM infiltrates
  - Sparse
  - interstitial
Immunophenotype

- CD2/3/5 and TCRβ positive
- CD4 positive in most cases
  - Elevated CD4/CD8 ratio
  - Increased proportion of CD4(+)/CD7(-) T cells
- CD8 expression is rare
- Aberrant T-cell phenotypes are common
Sezary Syndrome: Flow Cytometry
Genetics

- TCR clonally rearranged
- Complex karyotypes present in many patients
- No specific cytogenetic abnormality identified
Prognosis

- Aggressive disease
  - 10-20% 5 year survival rate

- May transform to a large T-cell lymphoma as a terminal event