

Spondyloarthritis Diseases

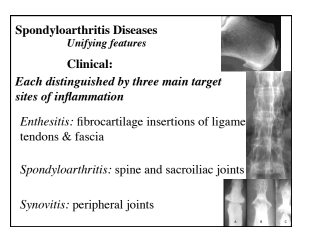
- Ankylosing spondylitis (ASp)
- ✓ Psoriatic arthritis (PsA)
- ✓ Reiter's syndrome (RS) / reactive arthritis (ReA)

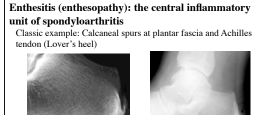
Undifferentiated spondyloarthritis (USpA)

Enteropathic arthritis (ulcerative colitis, regional enteritis)



Psoriasis, a related condition

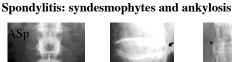




Features of inflammation:

•Infiltration of entheses by activated T cells

- •Granulation tissue forms (activated macrophages and fibroblasts)
- •Bone erosions and heterotopic new bone formation





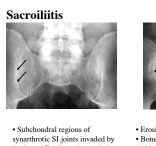
Activated T cells invade the junction of annulus fibrosis and vertebral body, triggering granulation tissue response



Annulus fibers eroded, the replaced by fibrocartilage: •Subperiosteal new bone formation •Fibrocartilage ossifies to form syndesmophytes

Inflammation resolves, b progressive cartilaginous and periosteal ossification

forms a "bamboo spine"

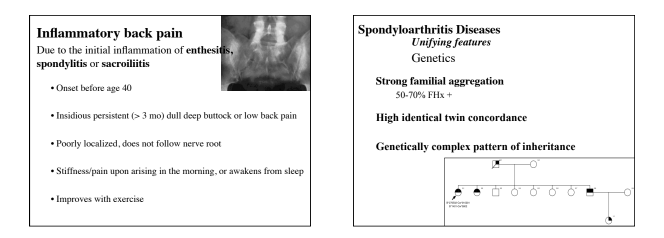


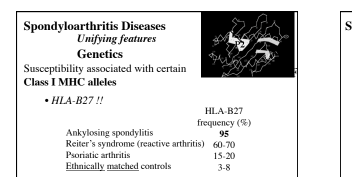
synarthrotic SI joints invaded by Activated T cells and granulation tissue



Erosion of cartilage on iliac side
 Bone plate blurring, joint space
 widening" and reactive sclerosis
 Fibrous ankylosis replaced by bone
 obliterating SI joint

Resolution of inflammation by heterotopic bone formation





• Other class I alleles also involved

Spondyloarthritis Diseases Unifying features

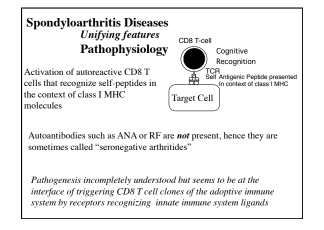
Pathophysiologic Mechanism

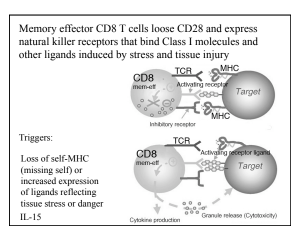
A clue from clinical medicine

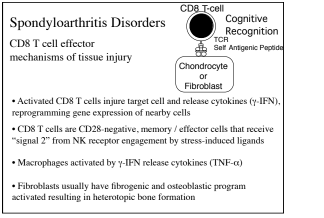
Unlike other autoimmune diseases that regress during development of AIDS, most spondyloarthritis diseases worsen or develop *de novo* at this time

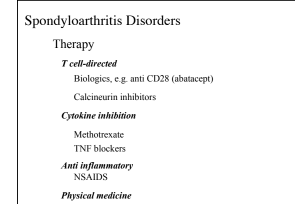
Implication:

CD4 T cells not required for development of symptomatic disease









Spondyloarthritis Diseases

- ✓ Ankylosing spondylitis (ASp)
- ✓ Psoriatic arthritis (PsA)
- ✓ Reiter's syndrome (RS) / reactive arthritis (ReA)



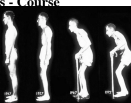
Ankylosing spondylitis

- · Widespread spondylitis and sacroiliitis
- Male: female =3-10:1
- Culminates in boney ankylosis of spine
- Onset, age 10-25 with dull pain in lumbar or gluteal regions
- Hip, shoulder knee arthritis in ~30%
- Epidemiology: >95% of those affected are HLA-B27
- Disease prevalence follows circumpolar distribution of HLA-B27
- Affects 1-3% of HLA-B27 individuals,
- No evidence for triggering by microorganisms

Ankylosing spondylitis - Course

Begins with sacroiliitis

 Inflammatory back pain and tenderness worsens and over sever months to years ascends, with increasing stiffness and loss of mobility



Postural changes: loss of lumbar lordosis, buttock atrophy and kyphosis; chest expansion compromised

• Peripheral joints, notably hips develop flexion contractures or ankylosis; compensatory knee flexion

 \bullet Peripheral arthritis (~30%) and peripheral enthesopathy (~30%) dominate the early phase of disease, then bony ankylosis predominates

Ankylosing spondylitis - systemic involvement

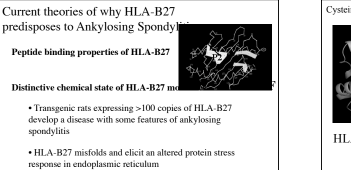
• Acute anterior uveitis (25%) may occur at any time; (syncheae and glaucoma)

- Apical pulmonary fibrosis, often with cavitation (<5%)
- Restrictive pulmonary disease due to costovertebral ankylosis (~ 10%)

 \bullet Granulomatous aortitis: complete heart block due to interventricular septum inflammation and /or aortic insufficiency (~5%)

Ankylosing spondylitis- different types of HLA-B27						
<i>HLA-B27</i> alleles differ from one another in polymorphic amino acids, in ethnic distribution and, importantly, whether they determine disease susceptibility						
Allele	Features	Ank.Spon				
B*2701	Rare	Yes				
B*2702	10% of AS in Europe and Middle East	Yes				
B*2703	Rare West African allele	Yes				
B*2704	Major HLA-B27 allele in China and India	Yes				
$\rightarrow B^{*2705}$	90% of AS, circumpolar Caucasians & Asia	ans Yes				
B*2706	SE Asia	No				
B*2707	Minor allele in SE Asia, China and India	Yes				
B*2708	Rare, UK and Azores	Yes				
B*2709	Sardinia	No				

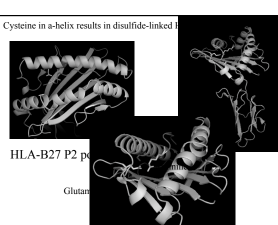
A self-peptide likely drives ankylosing spendulitie HLA-B27 alleles share the same P2 "B"pocke but differ from one another in the "F" P9 pock									
		P9 Pocket							
Allele	59	77	80	116	114	Ank.Spon			
B*2701	Tyr	Agn	Thr	Asp	His	Yes			
B*2702	Tyr	Agn	Ile	Asp	His	Yes			
B*2703	His	Asp	Thr	Asp	His	Yes			
B*2704	Tyr	Ser	Thr	Asp	His	Yes			
→ B*2705	Tyr	Asp	Thr	Asp	His	Yes			
B*2706	Tyr	Ser	Thr	Tyr	DAsp	No			
B*2707	Tyr	Asp	Thr	Asp	His	Yes			
B*2708	Tyr	Ser	Ile	Asp	His	Yes			
B*2709	Tyr	Asp	Thr	His	⊃His	No			



arthritis

% no

asis



Psoriasis / Psoriatic Arthritis

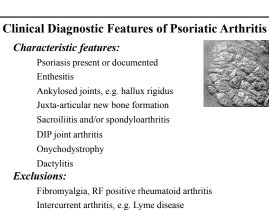
Psoriasis: skin disease with retardation in kertinocyte differentiation induced by activated T cells

Perhaps keratinocyte peptides are presented by class I molecules?

Psoriatic arthritis: spondloarthritis and psoriasis







Repetitive motion-induced musculoskeletal syndromes

Psoriatic arthritis - features

- Presentation: with obvious, subtle or no psoriasis, sometimes only isolated nail disease
- Onset typically insidious with stiffness; sometimes acute mimicking gout; can follow joint injury
- Sex: Male = female
- Early onset (<40 yrs) psoriatic arthritis has strong family history

Psoriatic arthritis

Dactylitis (Sausage digit) widespread inflammatory edema due to:

DIP and PIP arthritis of same ray Enthesitis

Tenosynovitis (flexor > extensor)

Periostitis

Onychodystrophy



Acral dystrophic state

Psoriatic arthritis

Enthesitis



•Sometimes subtle and easy to overlook

•Nonspecific foot pain, "tennis elbow" in the non dominant hand, or isolated posterior tibial tendinitis

•Widespread and symmetric, distribution differentiates from posttraumatic or occupational tendon injury

•Can be fulminant and combined with intense tenosynovitis

Psoriatic arthritis-peripheral joint patterns

Asymmetric oligoarthritis of small and medium-sized joints

Classic, with time more joints accumulate

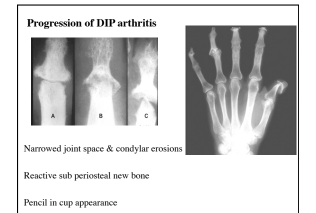
- DIP arthritis joints, also involves nails
 - Classic and unique to psoriatic arthritis, but only ${\sim}5{\text{-}}10\%$

Associated paronychia and swelling of the digital tuft may make appreciation of arthritis difficult; DDx Heberden's nodes

Arthritis mutilans

Osteolytic dissolution of joint with redundant overlying skin and telescoping digits (opera-glass hand)

Typical but uncommon; males and early-onset disease



Psoriatic arthritis-peripheral synovitis patterns

• Symmetric polyarthritis

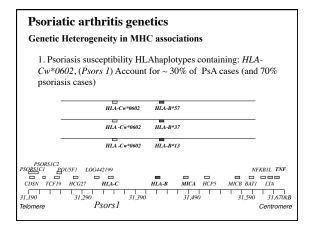
Most common pattern at onset, but is least specific for PsA

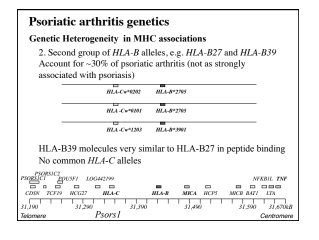
Hands, wrists, ankles, and feet

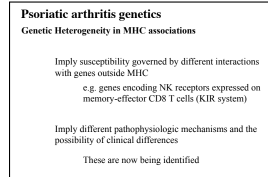
Differentiated from RA by enthesopathy and dactylitis, DIP joint involvement, relative asymmetry, new bone formation, pencil in cup deformity, absence of subcutaneous nodules, and negative RF

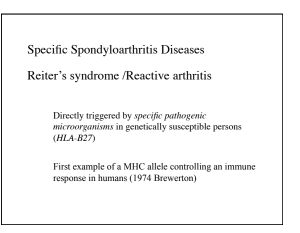
Important to distinguish RA from PsA because steroids contraindicated

Psoriatic Arthritis Genetics Psoriatic Arthritis-Nail Invo ~80-85% PsA, vs. 20-30% in Ps 6 Ø -Ø Nail matrix abnormalities ~60% strongly positive family Pitting histories, most often first degree · Onychodystrophy, crumbling relatives affected by psoriasis • Transverse ridging (Beau's lines) · Subungual hyperkeratosis Л · Leukonychia Onycholysis $\lambda_{\rm R}$ = 55 (assuming Ć · Ectatic capillaries prevalence 0.1%) Acral dystrophy · Nail matrix abnormalities Acrokeratosis Mode: mixed multifactorial pattern, partially dominant, incompletely · Often seen in digit involved with DIP arthritis penetrant









Reiter's syndrome /Reactive arthritis

"On August 21, 1916 a lieutenant in the Prussian army developed abdominal pain and diarrhea. This episode last 48 hours and was followed by a latent period of 7 days at which time urethritis and conjunctivitis occurred.

"The following day he developed polyarthralgias and arthritis of the knees, ankles, elbows, wrists and several interphalangeal joints.

"Within a few days the symptoms remitted and the patient remained well for 3 weeks.

"A relapse followed with a recurrence of urethritis and uveitis"

H. Reiter (Andre Calin)

Reiter's syndrome-clinical features I

• Onset 7- 30 days after specific enteric or venereal infection

• Course-Initial episode completely regresses, occasionally returns as increasingly intense recrudescences becoming chronic

 Peripheral arthritis: acute, highly inflammatory asymmetric arthritis involving knees, ankles, toes, and fingers (2-4 joints)
 All joints synchronous in abrupt fulminant onset

• Enthesitis - notably plantar fascia and Achilles tendon (40%)

• Dactylitis (Sausage digit) (40%)

· Sacroiliitis, stuttering spondyloarthritis

Reiter's syndrome Spondyloarthritis

Triad of Reiter's syndrome

Sub periosteal new bone formation a major feature



Infiltration of T cells

Fluffy reactive new bone formation

"Square" vertebrae but minimal paravertebral ossification

Asymmetric involvement of only one or two vertebral units

Reiter's syndrome - Clinical features II

 Onychodystrophy: subungual hype and para-keratosis

• Conjunctivitis (often first manifestation). Uveitis in recurrent disease

Non specific urethritis

• Painless circinate balanitis and mucosal ulcers, prostatitis

Heart - 10% of chronic phase 1° heart block from IV septum inflammation;
Aortic valve insufficiency due to granulomatous aortitis at aortic ring, rarely aortic dissection

Reiter's syndrome- role of specific infection

Develops 7-30 days after enteric infection with certain Gram neg. rods

- •Salmonella typhimurium, and occasionally S. paratyphi or S. heidelbergii
- •Shigella flexneri 2a and 2b, but not S. sonnei
- •Yersinea enterocoliticas
- •Campylobacter jejuni or C. fetus

These organisms typically invade and kill intestinal M cells, perhaps arthritogenic peptides cross-presented in class I MHC

Develops 7-30 days after venereal infection with

•*Chlamydia trachomatis* or *C. psittaci* Obligate intracellular eubacteria

Psoriasis / Reiter's syndrome in the setting of AIDS

Provided major clue pointing to importance of CD8 T cells in pathogenesis

Major source of disability in otherwise relatively well HIV+ patients in developing countries where HIV therapy is inadequate

