Disease Modifying Antirheumatic drugs





At the end of the lecture the students should: Know the pathogenesis of rheumatoid joint damage Emphasize the rational for early treatment of RA **Define and classify DMARDs** Compare and contrast the advantages and disadvantages of NSAIDs, Steroids and DMARDs in treatment of RA Know some examples of drugs related to DMARDs.

- Explore the pharmacokinetic and pharmacodynamic aspects of the selected DMARDs
- Describe the mechanism of action , specific clinical uses , adverse effects of individual drugs.

EPIDEMIOLOGÝ OF RHEUMATOID ARTHRITIS

Affects 1-2 % of the adult population

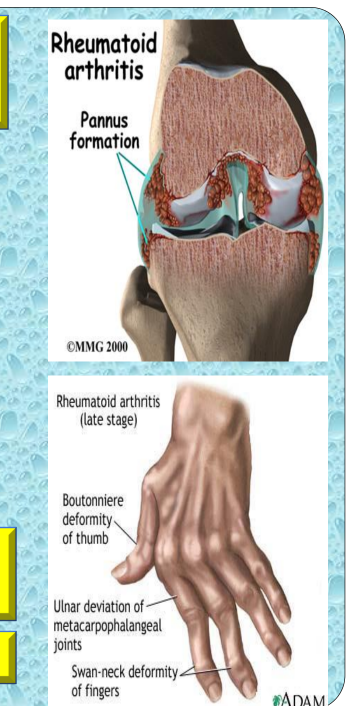
Is more common among women than in men (2-3 times)

Usually appears between ages 25 and 40 years

The incidence also increases with age, peaking between the 4th and 6th decades

Causes pain, disability and loss of function

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RHEUMATOID ARTHRITIS

RA is a chronic **autoimmune** disorder in which the normal immune response is directed against an individual's own tissue leading to:-

Decline in functional status

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Work disability &

socioeconomic costs

Systemic complications

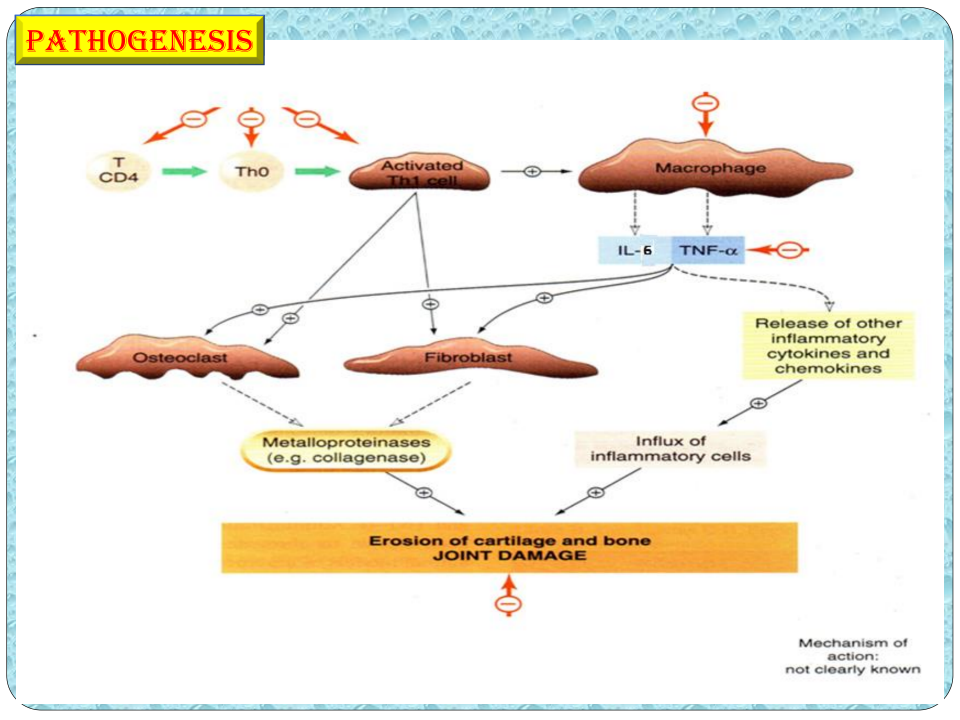
Co-morbidity & Increased mortality

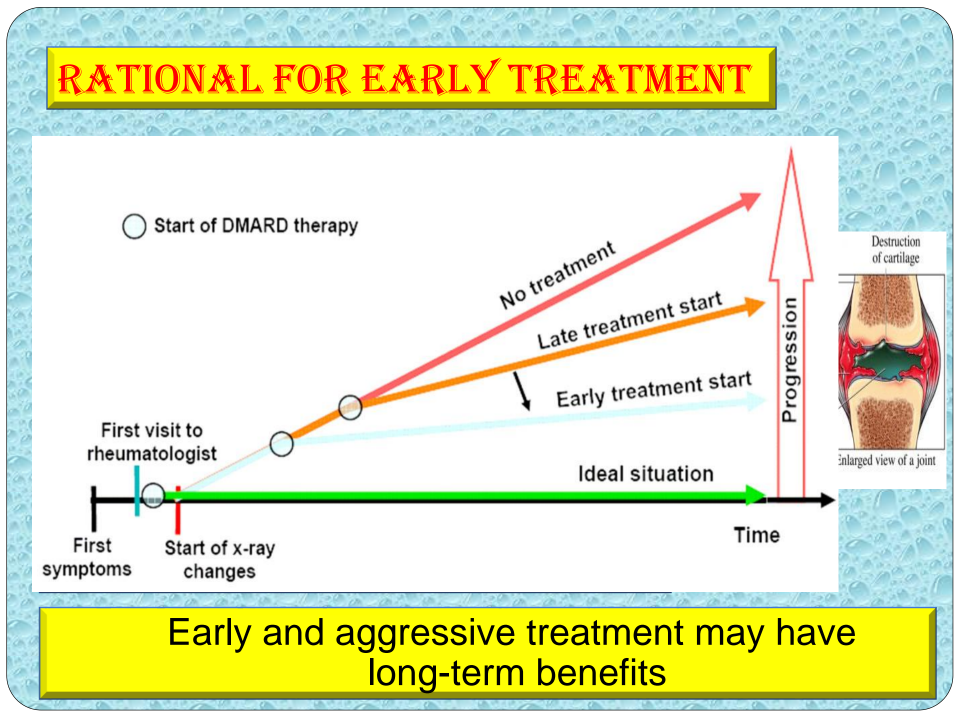


Synovial inflammation and hyperplasia, autoantibody production (rheumatoid factor), cartilage and bone destruction ("deformity")

> Anaemia Weight loss Osteoporosis Amyloidosis Renal, cardiovascular and neurological complications

RA damages joints and may be systemic





CLASSIFICATION

Drugs for Rheumatoid Arthritis

DMARDs

Biologic

Classical

NSAIDs

Glucocorticoids

GLUCOCORTICOIDS

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Anti-inflammatory drugs with an intermediate rate of action (slower than NSAIDs but faster than other DMARDs).

May be administered in low to moderate doses to achieve rapid disease control before the onset of fully effective DMARD therapy

Reserved for temporary control of severe exacerbations and long-term use in patients with severe disease not controlled by other agents.

Corticosteroids are too toxic for routine chronic use

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NSAIDS

Rapid onset of action

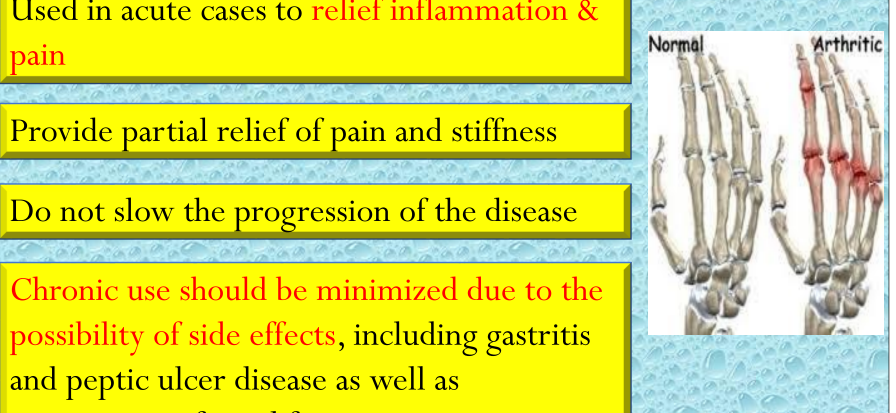
Used in acute cases to relief inflammation & pain

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Provide partial relief of pain and stiffness

Do not slow the progression of the disease

Chronic use should be minimized due to the possibility of side effects, including gastritis and peptic ulcer disease as well as impairment of renal function.



Comparison between NSAIDs & DMARDs

DMARDs

- Slow onset of action
- Arrest progression of the disease
- Prevent formation of new deformity
- Used in chronic cases when deformity is existing

NSAIDs

- Rapid onset of action
- No effect
- Can not stop formation of new deformity
- Used in acute cases to relief inflammation & pain

CLASSIFICATION OF DMARDS

Rheumatoid Arthritis

DMARDs act on the immune system to slow the progression of RA

Swollen inflamed Synovial membrane

Biologic

Infliximab Tocilizumab

Classical

Methotrexate Hydroxychloroquine

Bone erosion

GENERAL FEATURES

Used when the disease is progressing & causing deformities

Can not repair the existing damage, but prevent further deformity

Have no analgesic effects

Their effects take from **6 weeks up to 6 months** to be evident



METHOTREXATE

"Gold standard" for DMARD therapy & is the first-line DMARD for treating RA and is used in 50–70% of patients

Active in RA at much lower doses than those needed in cancer chemotherapy

MECHANISM

Inhibits dihydrofolate reductase

Reduces thymidine & purine synthesis

But at the dosages used for the treatment of RA, methotrexate has been shown **to stimulate adenosine release from cells,** producing an anti-inflammatory effect.

Inhibition of polymorphonuclear chemotaxis

Inhibition of T-Cells (cell-mediated immune reactions)

METHOTREXATE

PHARMACOKINETICS

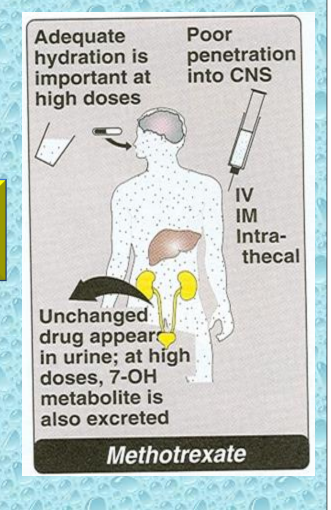
Approximately 70% absorbed after oral administration

Metabolized to a less active hydroxylated product

Half-life is usually only 6–9 hours

Excreted principally in the urine, but up to 30% may be excreted in bile.

Given 7.5 – 30 mg weekly





Bone marrow suppression

Dyspepsia, Mucosal ulcers

Hepatotoxicity

Pneumonitis

Teratogenicity

Leukopenia, anemia, stomatitis, GI ulcerations, and alopecia are probably the result of inhibiting cellular proliferation

Folic acid reduces GI & bone marrow effects

Monitoring:-Full blood count, ALT, Creatinine

HYDROXYCHLOROQUINE

MECHANISM

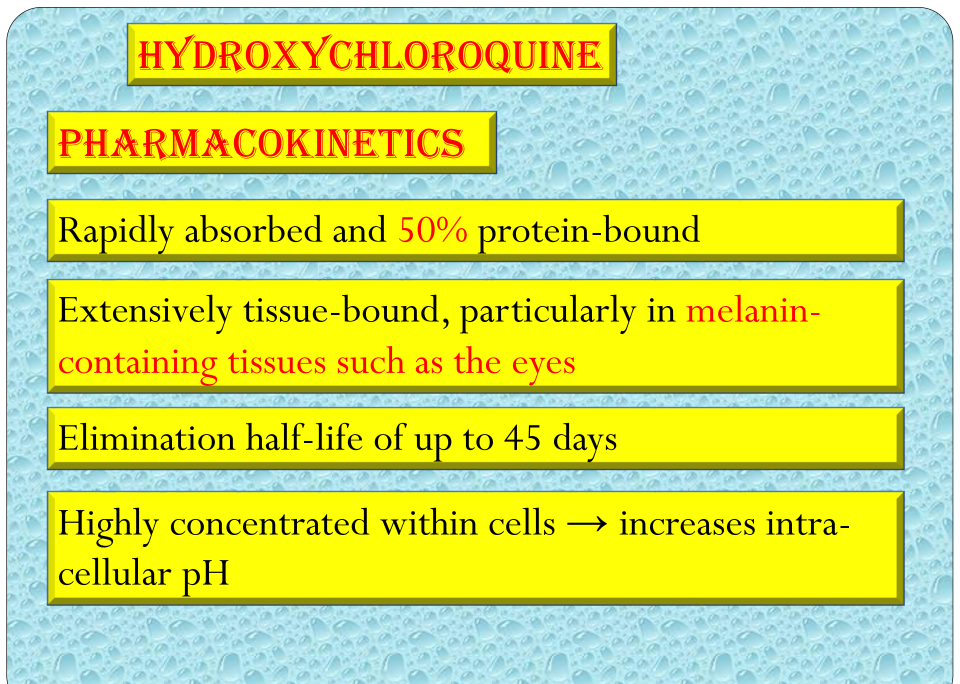
Stabilization of lysosomal enzyme activity

Trapping free radicals

Suppression of T lymphocyte cells response to mitogens

Inhibition of leukocyte chemotaxis





HYDROXYCHLOROQUINE

CLINICAL USES

Has not been shown to delay radiographic progression of disease

Generally used for treatment of early, mild disease or as adjunctive therapy in combination with other **DMARDs**.

Used in increasing methotrexate efficacy

Norma

Rheumatoid Arthritis

6 month response, mild anti-rheumatic effect

Rheumatoid - Hand





Least toxic, no blood tests is required

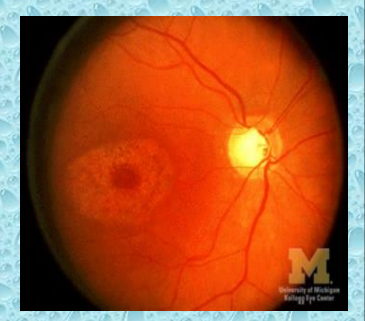
Nausea & vomiting

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Corneal deposits

Irreversible retinal damage



Ophthalmologic evaluation every 6 months

BIOLOGIC DISEASE MODIFIER

Genetically engineered drugs that are used to modify imbalances of the immune system in autoimmune diseases.

Some of these agents block, or modify the activity of selected cells in the immune system

They are expensive

Others work by blocking <mark>cytokines,</mark> that send signals between those cells

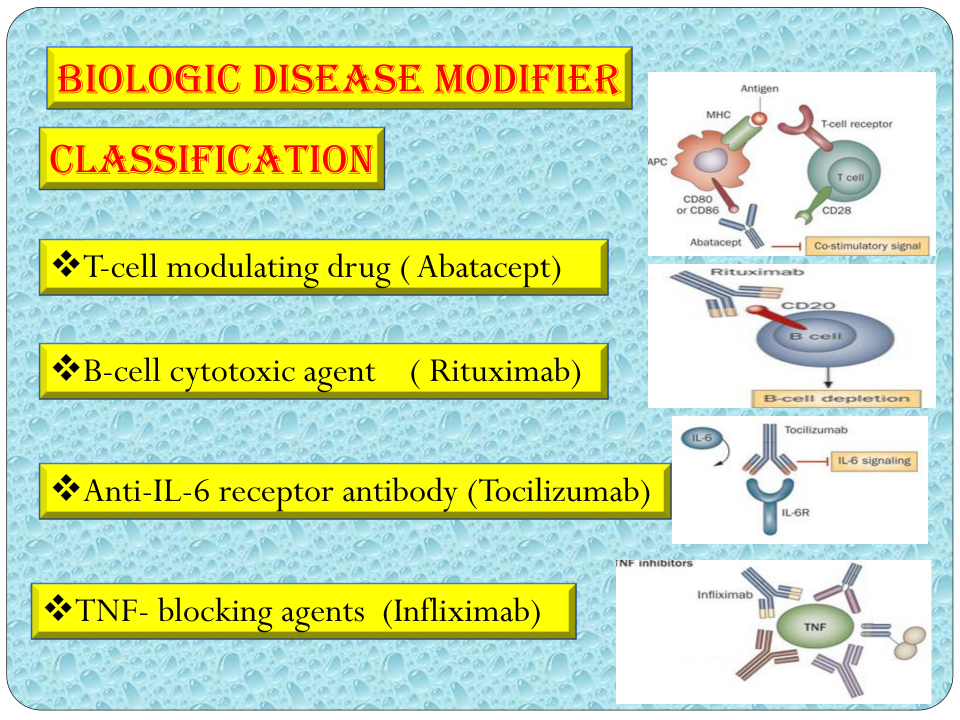
Abatacept Selectively Modulates T Cell Activation

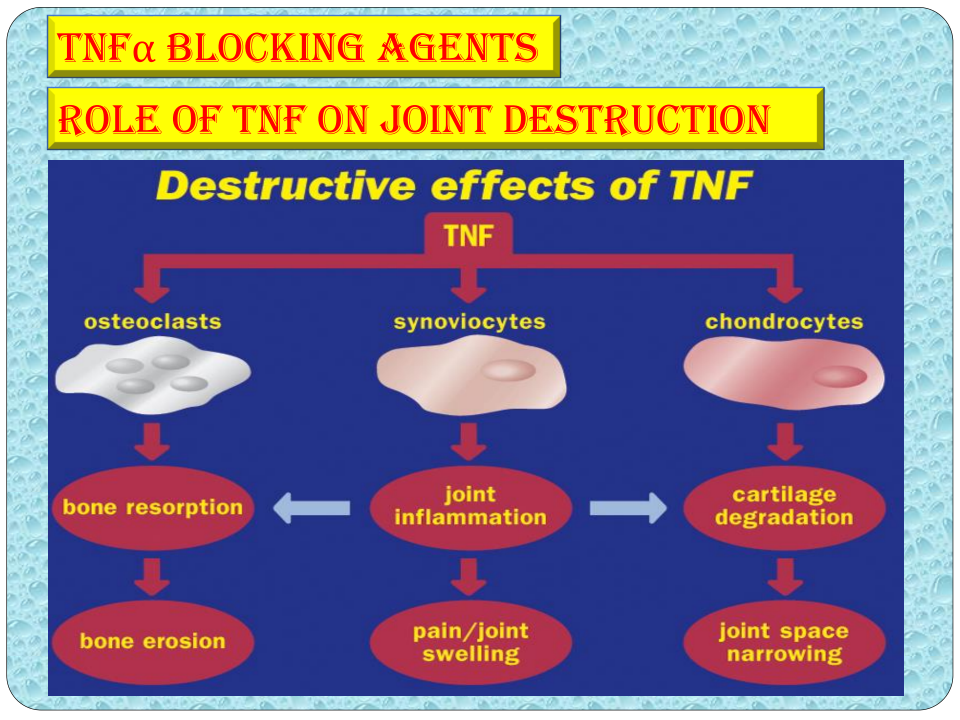
 Without Abatacept
 With Abatacept

 With Abatacept
 With Abatacept

 DC C D80/86 CD28
 CD20

 Activated T cell
 Abatacept





TNFα BLOCKING & GENTS

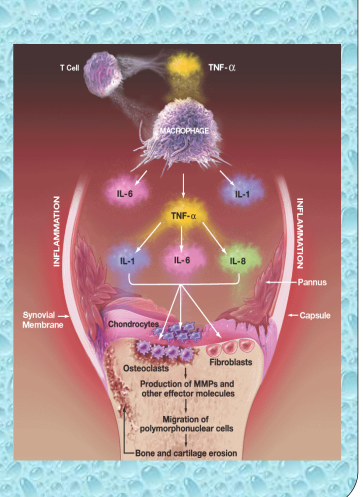
INFLIXIMAB

A chimeric IgG₁ monoclonal antibody (25% mouse, 75% human)

MECHANISM

It complexes with soluble TNF- α (and possibly membrane- bound TNF- α)and prevents its interaction with the cell surface receptors

This results in down-regulation of macrophage and T-cell function.



Monoclonal Antibody directed against TNF-Expline FAN SoR deprois Records a Enance dot alpha: Infliximab (Remicade B), Adalimumab (FINFel®)

SIGNAL

TNF

INFLIXIMAB

PHARMACOKINETICS

Given as an intravenous infusion with "induction" at 0, 2, and 6 weeks and maintenance every 8 weeks thereafter,

Terminal half-life is 9–12 days

After intermittent administration elicits human antichimeric antibodies in up to 62% of patients

Concurrent therapy with **methotrexate** decreases the prevalence of human antichimeric antibodies



INFLIXIMAB

CLINICAL USES

Infliximab is approved for use in RA, Ankylosing spnodilytis, Crohn's disease, ulcerative colitis,

It could be combined with methotrexate, hydroxychloroquine and other non biological DMARDs



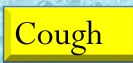
ADRS

Upper respiratory tract infections

Activation of latent tuberculosis

Infusion site reaction





Increase the risk of skin cancers including melanoma

TOCILIZUMAB

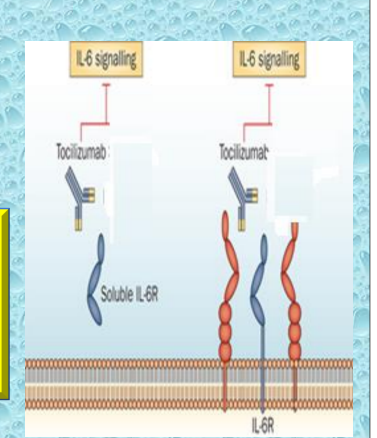
IL-6 is a proinflammatory cytokine implicated in the pathogenesis of RA,

With detrimental effects on both joint inflammation and cartilage damage

Tocilizumab binds to membrane IL-6 receptors, **blocking the activity of IL-6** in mediating signals that affect cytokine production, osteoclast activation

Half-life is dose-dependent

Given as monthly IV



TOCILIZUMAB

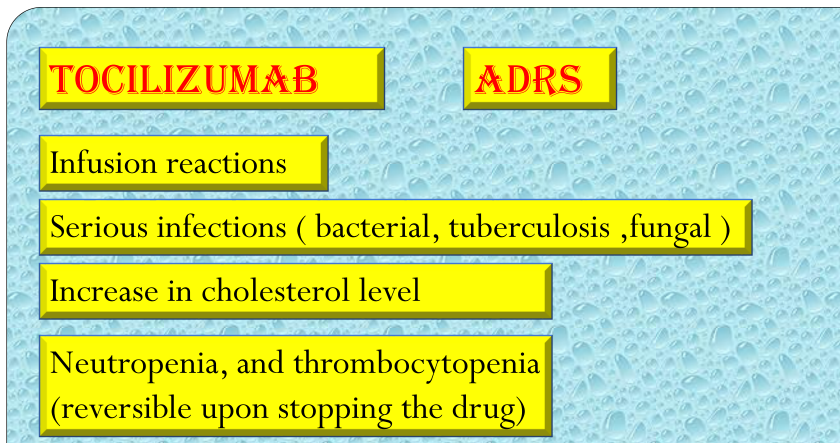
CLINICAL USES

Used as monotherapy in adult with rheumatoid arthritis or in children over 2 years with systemic juvenile arthritis

In combination with **methotrexate** or other non biologic anti-rheumatic drugs in patients with active rheumatoid arthritis not responding to TNF blockers or other biologic drugs







Decrease in WBCs

Increase in liver enzymes

Blood tests will be used monthly for increase in cholesterol, liver enzymes & decrease in WBCs

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TOCILIZUMAB

DRUG INTERACTIONS

IL-6 inhibits CYP450

Tocilizumab restores the activity of the enzyme (essential for the metabolism of some drugs such as cyclosporine, warfarin).

BEST WISHES

