

Rheumatic Heart Disease

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Objectives

- To understand basis of rheumatic fever as an immunologically mediated **late complication** of Streptococcal infection
- To know that autoimmunity results from production of **cross reacting antibodies** against Streptococcal antigens
- To describe rheumatic **heart disease** as one of the several manifestations of rheumatic fever
- To know the signs, symptoms, pathogenesis, treatment and prophylaxis of rheumatic heart disease

Rheumatic Fever

- Epidemiology of Rheumatic Fever (RF)
- ~3% of persons with untreated group A streptococcal **pharyngitis** develop rheumatic fever
- 15-20 million new cases a year in developing countries
- **Risk factors**
 - Low standard of living
 - Crowding

Rheumatic fever

- Individual (HLA) susceptibility is also important
- Antigen-presenting cells bearing the **HLA-DR7** molecule from RHD patients preferentially recognize heart-tissue protein

(Guilherme L, Kalil J. Ann N Y Acad Sci 2007,1107:426-433)

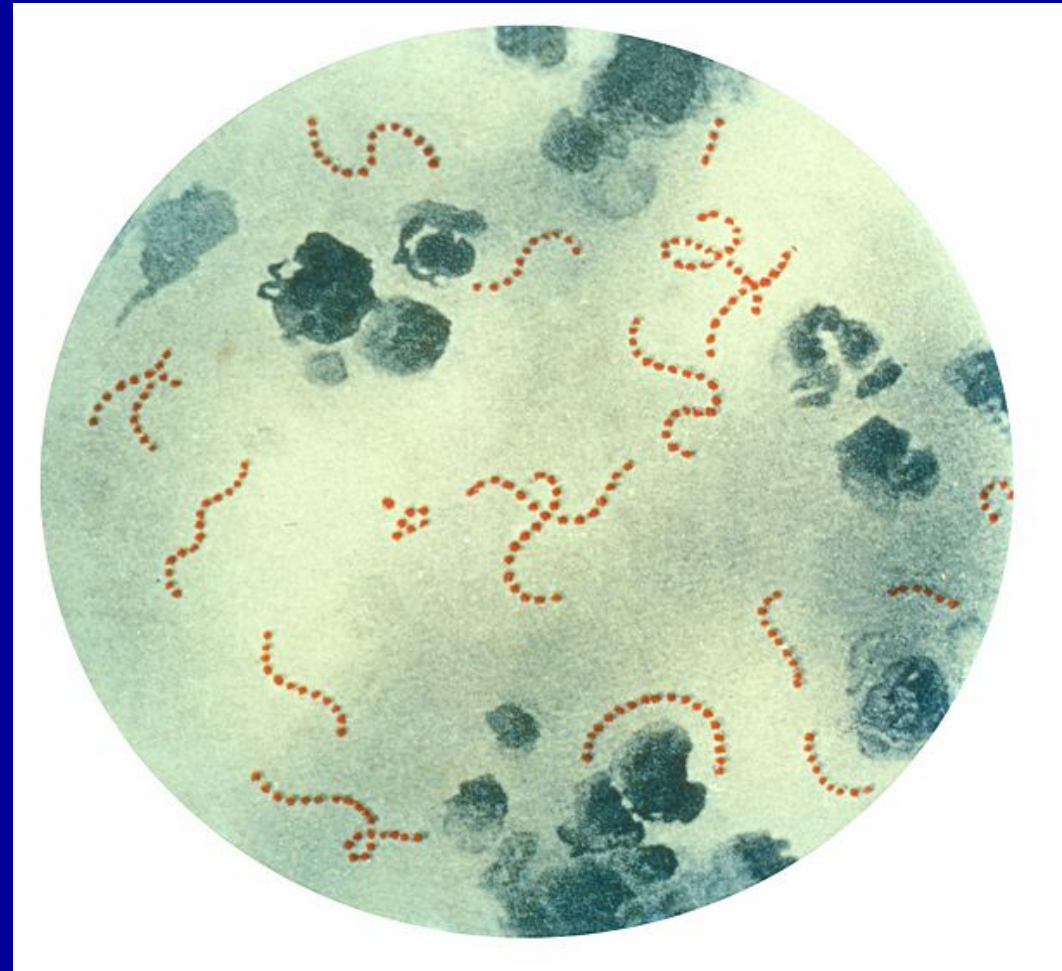
- Other views in the literature exist, due to
 - + The various HLA-typing methods.
 - + Ways of grouping the cases.

Rheumatic fever

- **Rheumatic fever** is an inflammatory disease which may develop after a **Group A Streptococcal infection** such as:
 - Strep. throat infection or scarlet fever
- Can involve the **heart, joints, skin, and brain**
- It commonly appears in children ages **5 through 15**

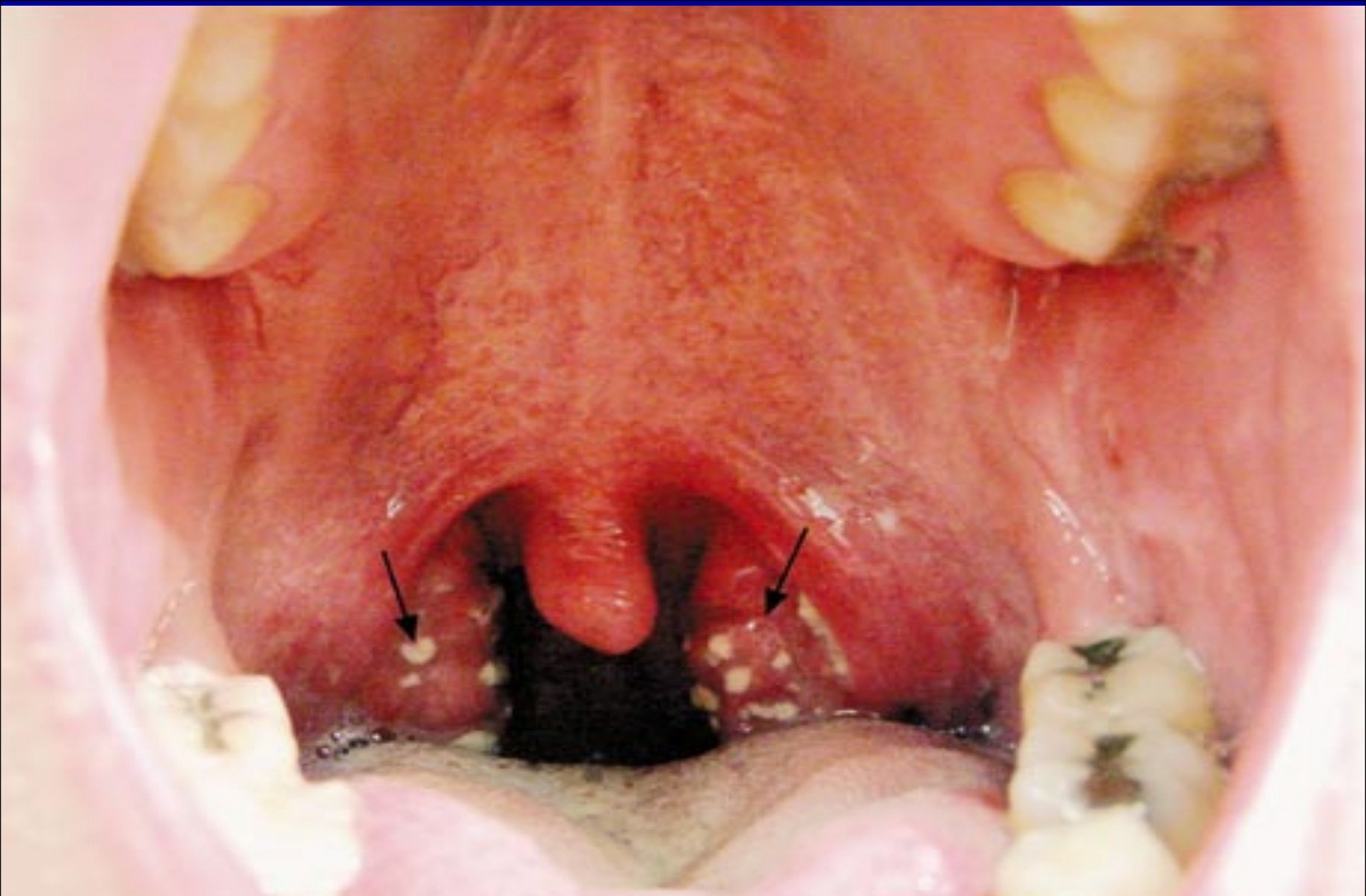
Organism

- Caused by **group A streptococcus**
- There is a latent period of **~3 weeks (1–5 weeks)** between the group A streptococcal infection and the appearance of the clinical features of RF



Group A β -haemolytic streptococcus

- All cases associated with recent infection (e.g. pharyngitis)
- **Antibody and cellular immune response** cross-reacts with human connective tissue



**Nimishikavi S, Stead L Streptococcal Pharyngitis – Images in Clinical Medicine.
NEJM 2005; 352:e10.**

M proteins

Adherence of *Streptococcus pyogenes* to host cells & inhibiting the host immune response

Hyaluronic acid capsule: Camouflages the bacterium

Streptokinases: Dissolve blood clots

Peptidases:

Degrades proteins involved in immune response

Pyrogenic toxins: Stimulate fever, rash & shock

Streptolysins:

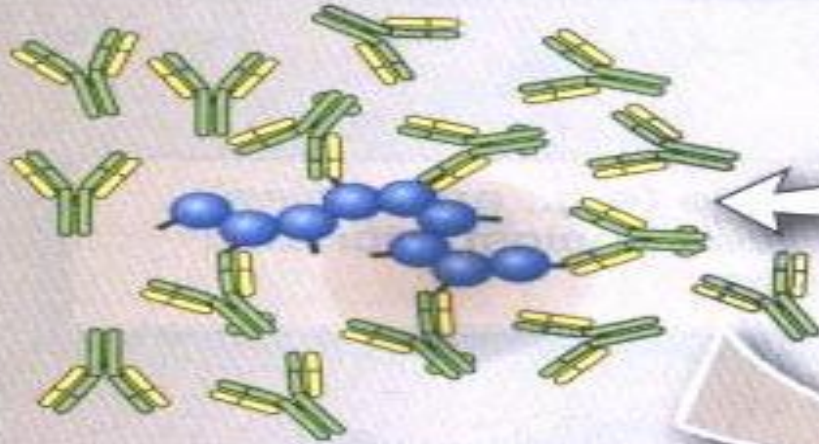
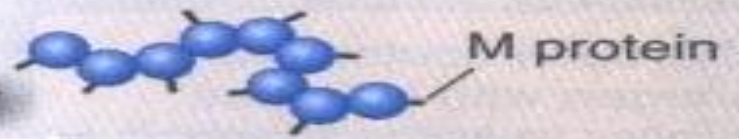
Lyse erythrocytes, leukocytes & platelets

PATHOGENESIS

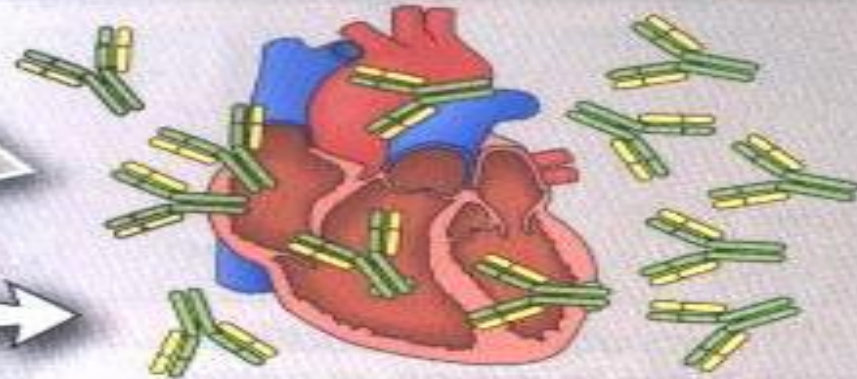
- Rheumatic fever affect the peri-arteriolar connective tissue
- It is believed to be caused by **antibody cross-reactivity**
- This cross-reactivity is a **Type II hypersensitivity** reaction and is termed *molecular mimicry*

- Group A streptococcus pyogenes has a cell wall composed of branched polymers which sometimes contain "**M proteins**" that are highly antigenic
- The **antibodies** which the immune system generates against the "**M proteins**" may cross react with **cardiac myofiber protein myosin and smooth muscle cells of arteries**, inducing cytokine release and tissue destruction
- This inflammation occurs through direct attachment of **complement** and **Fc receptor-mediated recruitment of neutrophils and macrophages**

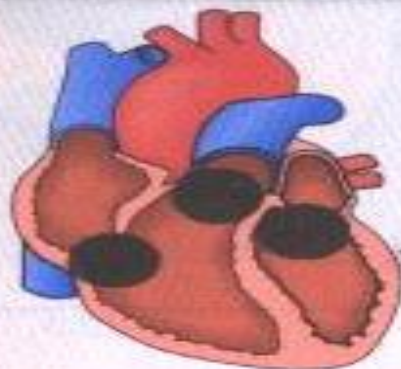
Group A Strep present in throat infection; expresses large amounts of M protein



Large amounts of IgG produced against M protein on *Strep*



Antibodies against M protein can bind to molecules on cardiac cells that are very similar to M protein



Antibody-induced injury to heart valves and sarcolemma

Rheumatic Valve Model

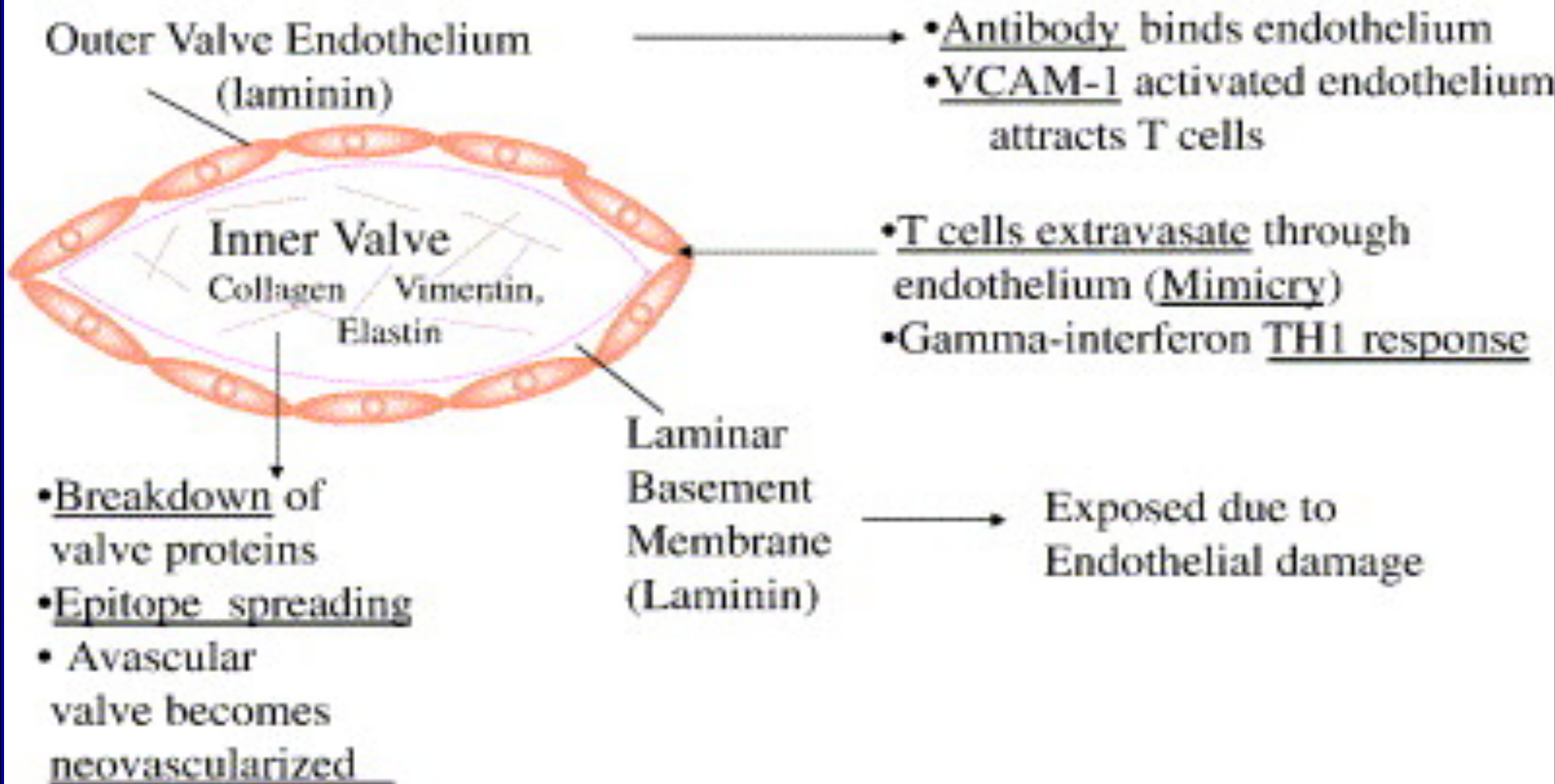
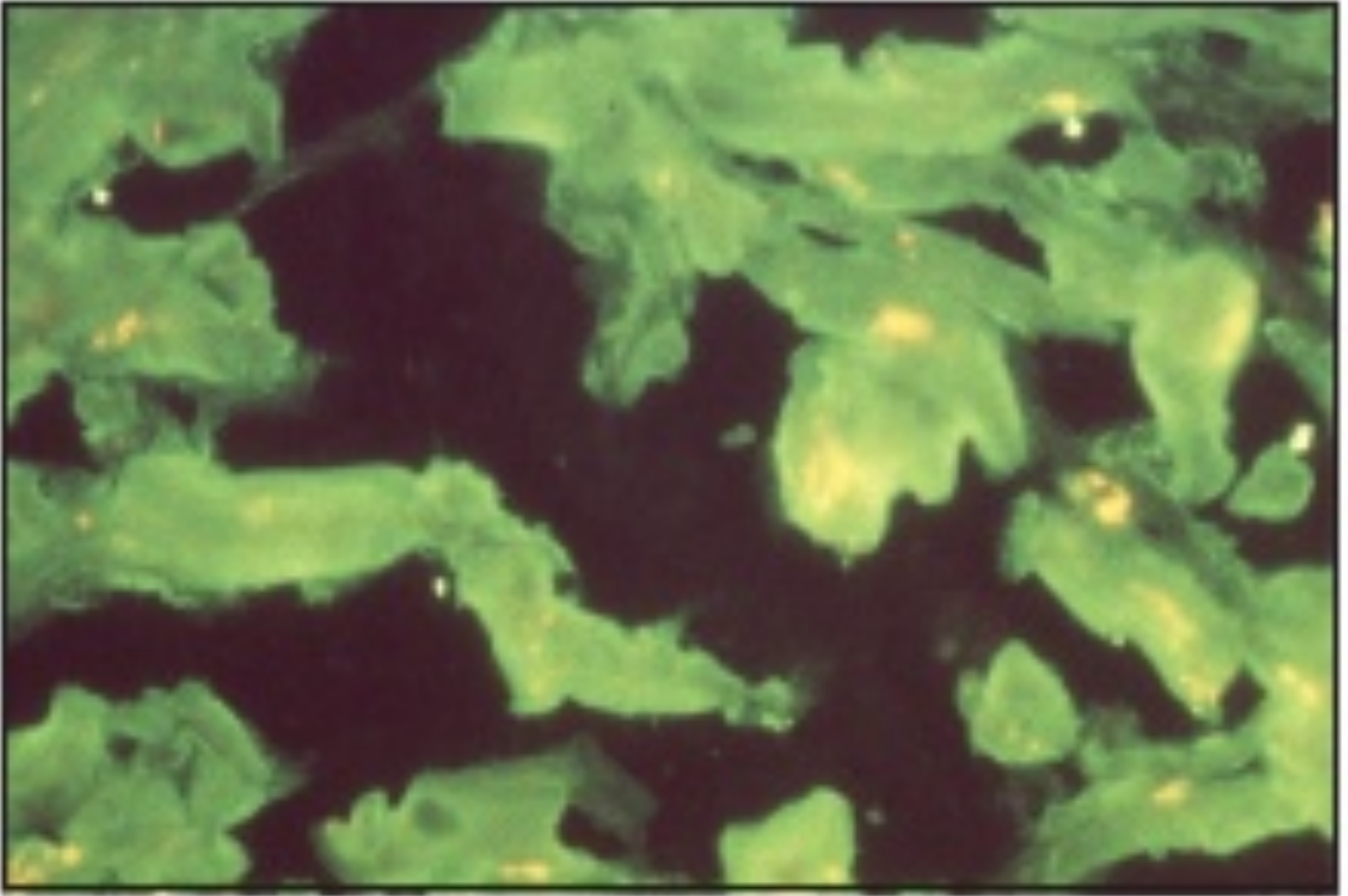


Diagram illustrating the process of initial mimicry which leads to **granuloma formation, gamma interferon production and scarring in the valve**. After the initial process has developed inflammation in the valve, other proteins in the valve may then be recognized by the immune system leading potentially to **epitope spreading** and responses against other valve proteins such as **vimentin and collagen**.

"Molecular mimicry in the autoimmune pathogenesis of rheumatic heart disease" by L. Guilherme; J. Kalil; M.W. Cunningham.



Immunofluorescent staining of heart muscle with serum obtained from an acute rheumatic fever patients

Pathophysiology

- During a Strep. infection activated antigen presenting cells such as macrophages present the bacterial antigen to helper T cells
- Helper T cells subsequently activate self reactive B cells and induce the production of antibodies against the cell wall of Streptococcus
- However the antibodies may also react against the myocardium and joints, producing the symptoms of rheumatic fever

Environmental factors, especially overcrowding

Precipitating event: infection with a strain of group A streptococcus carrying specific virulence factors

Repeated group A streptococcus infections

Repeated or ongoing infections possibly driving the valvular inflammatory response

Susceptible host

Priming of immune response

First episode of ARF

Episodes of recurrent ARF

RHD

Molecular mimicry between group A streptococcus antigens and host tissues

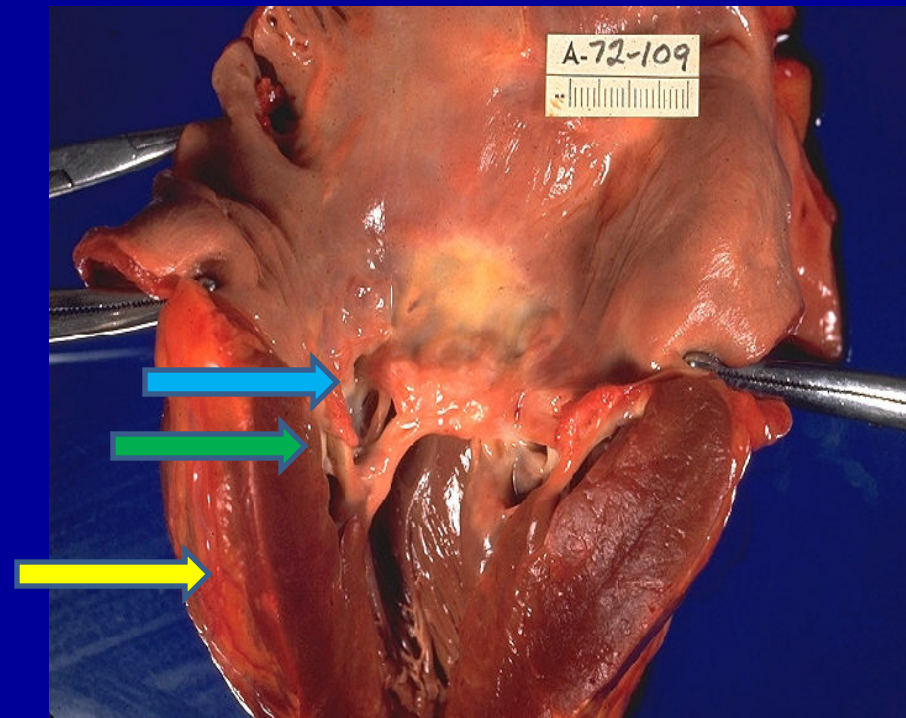
Exaggerated T-cell mediated immune response

Genetically determined host factors

Clinical Presentation

Heart

- Up to 60% of patients with ARF progress to Rheumatic Heart Disease (RHD)
- **only manifestation of ARF with significant potential to cause long-term disability and/or death**
- The endocardium, pericardium, or myocardium may be affected (**pancarditis**)
- Valvular damage is the hallmark of rheumatic carditis. The **mitral valve** is almost affected



Left ventricle has been cut open to display characteristic severe thickening of mitral valve, thickened chordae tendineae, and hypertrophied left ventricular

Joints (arthritis)

- This is usually polyarthritis, sometimes **flitting** from joint to joint (migratory), affecting the larger joints more than the smaller ones.
- Swelling, redness and tenderness are the common findings and occasionally joint effusions
- Inflamed **Keen Joint**



Skin (Erythema Marginatum)

- **Skin lesions:** The classical erythema marginatum—lesions with prominent margins slightly raised



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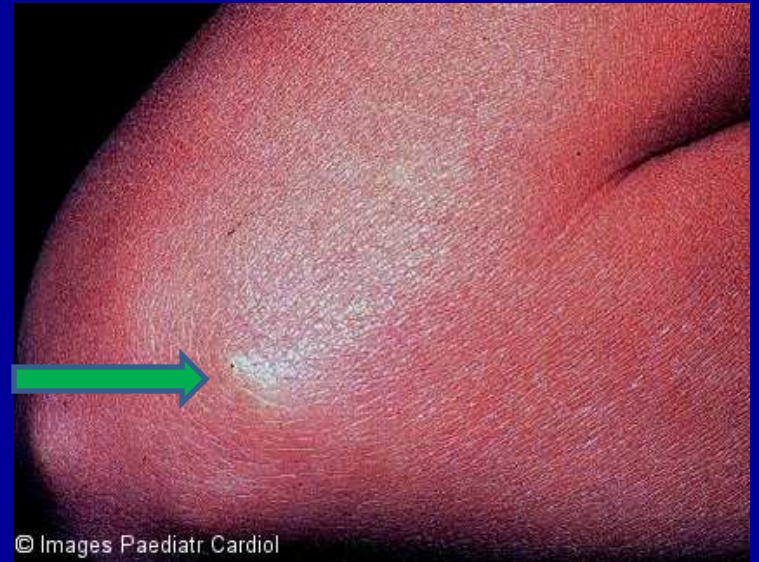
Central nervous system (chorea)

- Sydenham's chorea
- Occurs in children, rare in adults
- The choreiform movements affect particularly the head and the upper limbs
- They may be generalized or restricted to one side of the body (hemi-chorea)
- Chorea eventually resolves completely, usually within 6 weeks
- likely due to molecular mimicry, with autoantibodies reacting with brain ganglioside

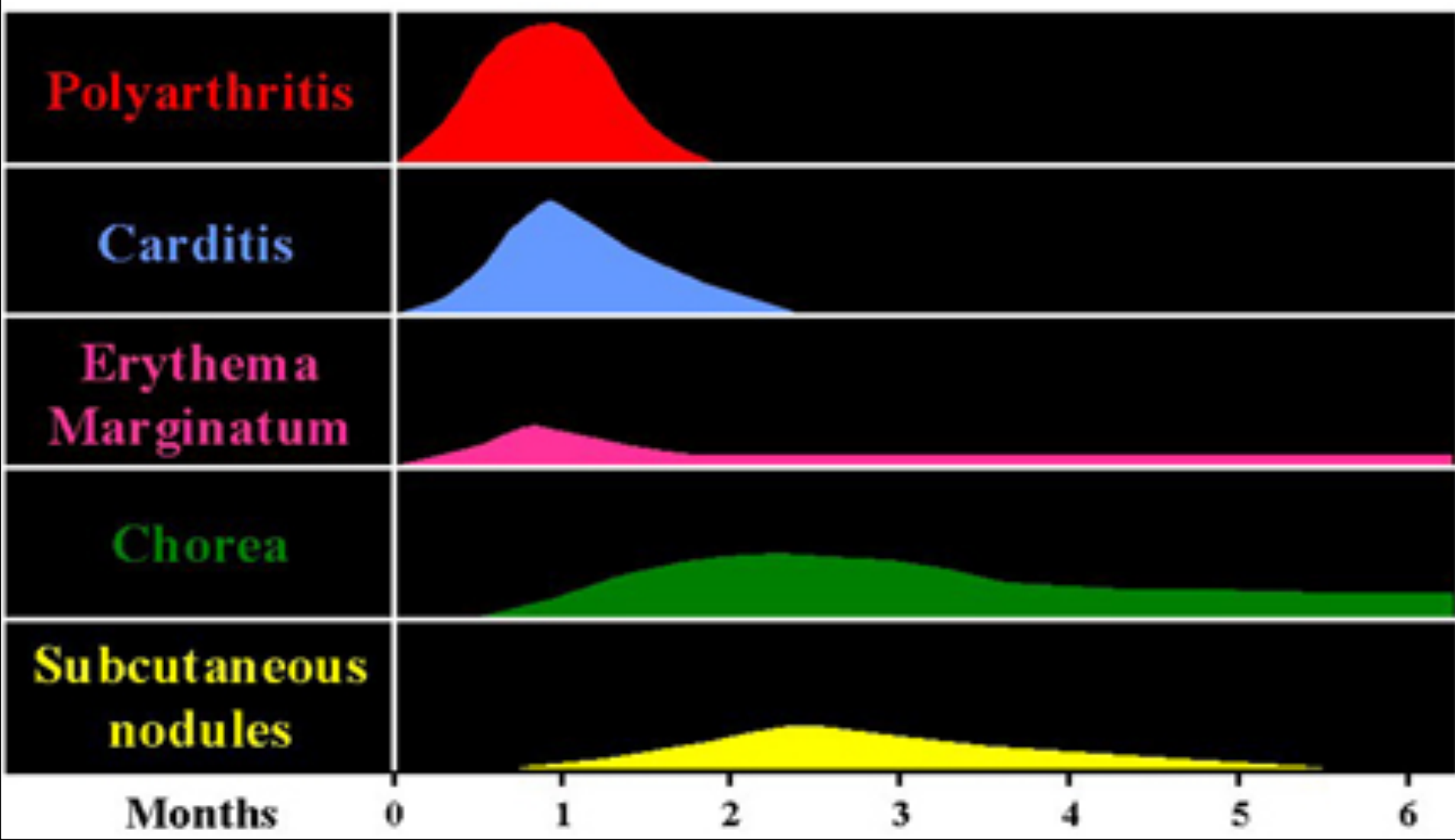


Subcutaneous nodules

- **Subcutaneous nodules :**
These are painless, round, firm lumps overlaid by normal looking skin
- They range from a few millimeters to 1.5 cm in diameter, and are localized over bony prominences like the elbow, shin and spine. They sometimes last longer than a month



Clinical Manifestations of Acute Rheumatic Fever



Investigation of Rheumatic Fever

- Anti-streptolysin O (ASO) titer
 - At least 80% of patients with ARF have an elevated **anti-streptolysin O** titer at presentation
 - Rising titer is more convincing
 - **Anti-DNAse B**
 - **Anti-hyaluronidase test**
- **Throat culture** for group A streptococci (obtain 2 or 3 cultures)

Rheumatic Fever – Clinical Course

- Subsequent attacks
 - Increased vulnerability to **reactivation of disease** with subsequent strep infections
 - **Same symptoms** with each attack
 - **Carditis** worsens with each attack
 - **Heart valves** are frequently deformed (mitral)
 - **Heart failure** develops after decades

Acute, recurring, chronic:

- Symptoms prone to recur with subsequent Strep. infections
- Chronic disease leads to fibrosis (chordae of heart valves + valve cusps)



Stenotic mitral valve seen from left atrium



Opened stenotic mitral valve

Treatment of Rheumatic Fever

- Treat first strep throat infection with **penicillin**
- Treat other manifestations **symptomatically**
- **Prophylactic** long term anti-strep therapy given to anyone who has had rheumatic fever

Take home message

- Rheumatic heart disease results from **cross reacting antibodies** binding the heart valves
- Repeated attacks of Streptococcal throat infection over the years damage heart valves resulting in either **stenotic or incompetent** heart valves
- Treatment involves **surgical** replacement of the damaged heart valves
- In patients with rheumatic fever long term administration of **penicillin** is recommended for prevention of future infections by group A Streptococcus

Thank you