Rheumatic Fever And RHD

Ayman Al-Saleh, MD, MSc, DABIM(C), FACP, FRCP(C) Interventional and Structural Cardiology

Assistant Professor (Adjunct), McMaster University. Hamilton, Canada.

Assistant Professor, King Saud University. Riyadh, Saudi Arabia

Lecture Outline

What is ARF And RHD?

Diagnosis

Jones Criteria

Differential Diagnosis

Investigations, Management

Rheumatic Valvular Heart Disease

Prevention

Rhuematic Fever

Follows group A beta hemolytic streptococcal throat infection

It represents a delayed immune response to infection with manifestations appearing after a period of 2-4 weeks

□Age 5-15 yrs

A multisystem disease

RHD is a long-term complication of ARF

•Major effect on health is due to damage to heart valves

Pathologic Lesions

 Ashcoff nodules
 Fibrinoid degeneration of connective tissue, inflammatory cells



Global Burden of RHD-WHO

A leading cause of CV morbidity & mortality in young people

□Total cases with RHD 15.6 Millions

CHF:3Million,valve surgery required in 1Million

Annual incidence of RF: 0.5 Million, nearly half develop carditis

Estimated deaths from RHD: 250,000/YR

Imposes a substantial burden on health care systems with limited budgets

Epidemiologic Background

The incidence of RF and the prevalence of RHD has declined substantially in Europe, North America and other developed nations

this decline has ben attributed to improved hygiene, reduced household crowding, and improved medical care

Epidemiologic Background

The major burden is currently found in low and middle income countries (India, middle east), and in selected indigenous populations of certain developed countries (Australia and Newzealand).

A disease of poverty and low socioeconomic status

In underdeveloped countries RHD is the leading cause of CV death during the first five decades of life

ARF and RHD in Saudi Arabia

- In developed countries the incidence of ARF has declined over past 50 years, incidence ranging 0.2 -0.64/100,000 (USA).
- ARF incidence in Eeastern province of SA was 22/100,000, age 5-14 years.
- ARF incidence in Kwait 29/100,000, age 5-14 years
- A study from Western province showed a prevalence of RHD 2.4/1000, age 6-15 years.
- A large study showed an overall prevalence of ARF and RHD in SA 3.1/1000 , age 6-15
- In Egypt prevalence of RHD, 5/1000 school-children
- In Yemen 36.5/1000 school-children

Diagnosis of ARF

□No single test to diagnose ARF

The symptoms and signs are shared by many inflammatory and infectious diseases

•Accurate diagnosis is important

Overdiagnosis will result in individuals receiving treatment unnecessarily

Underdiagnosis may lead to further episodes of ARF causing damage, and the need for valve surgery, and or premature death

Diagnosis of ARF

Diagnosis is primarily clinical and is based on a constellation of signs and symptoms, which were initially established as the Jones criteria

In 1944 Dr. TD Jones published a set of guidelines for diagnosis of ARF "Jones Criteria"

Subsequently Modified in 1965, 1984 and 1992by AHA

Revised recently -2015 by AHA

1992 Modified Jones Criteria

| Major criteria | Minor criteria |
|-------------------------|--|
| Migratory polyarthritis | Arthralgia |
| Carditis | Fever |
| Erythema marginatum | First degree heart block |
| Syndenham chorea | Elevated inflammatory markers (ESR, CRP) |
| Subcutaneous nodules | |

Carditis

□Occurs in 50-70% of cases

Only manifestation of ARF that leaves permanent damage

□May be subclinical

Murmurs of MR or AR may occur in acute stage while mitral stenosis occurs in late stages

Cardiomegaly and CHF may occur

Arthritis

Common: present in 35-66%

Earliest manifestation of ARF

Large joints: The knees and ankles, shoulders, elbows

• "Migrating", "Fleeting" polyarthritis

Duration short < 1 week

Rapid improvement with salicylates

Does not progress to chronic disease

Sydenham Chorea

DAlso known as Saint Vitus'dance

Occur in 10-30%, extrapyramidal manifestation, female predominnce

Abrupt Purposeless involuantry movements of muscles of face, neck, trunk, and limbs.

Delayed manifestation of ARF -months

Clinically manifest as-clumsiness, deterioration of handwriting, emotional lability or grimacing of face

Subcutaneous Nodules

- Occur in 10%
- \Box Usually 0.5 2 cm long
- □Firm non-tender
- Occur over extensor surfaces of joints, on bony prominences, tendons, spine
- Short lived: last for few days
- **Associated** with severe carditis

Subcutaneous Nodules





Erythema Marginatum

 \Box Present in <6%

Less common, but highly specific manifestation of ARF

Reddish border, pale center, round or irregular serpiginous borders, non-pruritic, transient rash

Occurs on trunk, abdomen or proximal limbs

Associated with carditis

Erythrma Marignatum



Revised Jones Criteria-2015

| A. For all patient populations with evidence of preceding GAS infe | ction |
|--|---|
| Diagnosis: initial ARF | 2 Major manifestations or 1 major plus 2 minor manifestations |
| Diagnosis: recurrent ARF | 2 Major or 1 major and 2 minor or 3 minor |
| B. Major criteria | |
| Low-risk populations* | Moderate- and high-risk populations |
| Carditis† • Clinical and/or subclinical | Carditis Clinical and/or subclinical |
| Arthritis • Polyarthritis only | Arthritis • Monoarthritis or polyarthritis • Polyarthralgia‡ |
| Chorea | Chorea |
| Erythema marginatum | Erythema marginatum |
| Subcutaneous nodules | Subcutaneous nodules |
| C. Minor criteria | |
| Low-risk populations* | Moderate- and high-risk populations |
| Polyarthralgia | Monoarthralgia |
| Fever (≥38.5°C) | Fever (≥38°C) |
| ESR \geq 60 mm in the first hour and/or CRP \geq 3.0 mg/dL§ | ESR \geq 30 mm/h and/or CRP \geq 3.0 mg/dL§ |
| Prolonged PR interval, after accounting for age variability (unless carditis is a major criterion) | Prolonged PR interval, after accounting for age variability (unless carditis is a major criterion) |

2015 Revised Jones Criteria

A firm diagnosis requires

1) 2 Major manifestations or 1 Major and 2 Minor manifestations

and

2) Evidence of a recent streptococcal infection.

2015 Revised Jones Criteria

Evidence of Preceding GAS Infection:

- 1) Increased or rising ASO titer or Anti-Dnase B titer
- 2) A positive throat culture

Rheumatic Fever Recurrences

- Reliable past history of ARF: 2 major or 1 major and 2 minor or 3 minor manifestations sufficient for diagnosis
- Presence of antecedent streptococcal infection
- When minor manifestations only present exclude other causes.

DDX of ARF

| Presentation | | | | |
|--|-----------------------------------|---|--|--|
| Polyarthritis and fever | Carditis | Chorea | | |
| Septic arthritis (including | Innocent murmur | Systemic lupus erythematosus | | |
| disseminated gonococcal infection)* | Mitral valve prolapse | Drug intoxication | | |
| Connective tissue and other autoimmune disease ⁺⁺ | Congenital heart disease | Wilson's disease | | |
| Viral arthropathy [¥] | Infective endocarditis | Tic disorder [±] | | |
| Reactive arthropathy [¥] | Hypertrophic cardiomyopathy | Choreoathetoid cerebral palsy | | |
| Lyme disease≠ | Myocarditis: viral or idiopathic | Encephalitis | | |
| Sickle cell anaemia | Pericarditis: viral or idiopathic | Familial chorea (including Huntington's) | | |
| Infective endocarditis | | Intracranial tumour | | |
| Leukaemia or lymphoma | | Lyme disease [≠] | | |
| Gout and pseudogout | | Hormonal [§] | | |

Investigations

Recommended for all cases

White blood cell count

Erythrocyte sedimentation rate (ESR)

C-reactive protein (CRP)

Blood cultures, if febrile

Electrocardiogram (if prolonged P-R interval or other rhythm abnormality, repeat in 2 weeks and again at 2 months, if still abnormal

Chest X-ray, if clinical or echocardiographic evidence of carditis

Echocardiogram (consider repeating after 1 month, if negative)

Throat swab (preferably before giving antibiotics): culture for group A streptococcus

Antistreptococcal serology: both ASO and anti-DNase B titres, if available (repeat 10–14 days later if first test not confirmatory)

Treatment of ARF

Bed rest

Salicylates : Aspirin

- •75-100 mg /kg/day given as 4 divided doses for 6 -8 weeks
- •Attain a blood level 20-30 mg/dl
- Penicillin: Procaine Penicillin 4 million units/day x10 days
- Prednisolone:2mg/kg/day taper over 6 weeks, Given when there is severe carditis

Heart Failure Treatment: diuretics, ACEI

Chronic Rheumatic Heart Disease

- □Most commonly in Mitral-70%
- **□**Frequently in Aortic-40%
- Less frequently Tricuspid-10%
- Rarely pulmonary valve-2%
- Mitral Stenosis is more common in females(3:1), while males have higher incidence of Aortic Regurgitation

Mitral Stenosis

The normal MVA= 4-5 cm2
In severe ms <1.5 cm2
High LAP

The rise in LAP causes a similar rise in pulmonary capillaries, veins and artery

Mitral Stenosis



Clinical Features

Dyspnea **Fatigue** Palpitation □Hemoptysis (10%) Hoarseness (Ortner's syndrome) Dysphagia □Storke or peripheral embolization

Clinical Features

Cyanosis (Mitral facies, malar flush) □Tapping apex (S1) □Parasternal heave Diastolic thrill □Accentuated S1, accentuated S2 Opening snap □Mid-diastolic rumble

Investigations

- Straightening of the left heart border
- Double density
- Kerley B lines , CA in MV
- ECG: LAE, P Mitrale ,RV dominance
- Echodoppler

Echo In Mitral Stenosis



Management

B-Blockers ,CCB
Digoxin (AF)
Warfarin
Balloon Valvuloplasty
Mitral valve replacement

BMV



Mitral Regurgitation

Asymptomatic Dyspnea, orthopnea, PND Displaced PMI, Thrill □Soft S1, □Pansystolic murmur □Treatment is surgical

ECHO



Aortic Regurgitation-Signs

□Water-hammer / collapsing pulse □Wide pulse pressure □Corrigan's sign De Musset sign □Muller sign Quincke's pulse □Hill's sign

ECHO



Aortic Stenosis



Symptoms

AnginaSyncopeDyspnea

Signs

- □Arterial Pulse wave form : Plateau
- Small (Parvus)
- Slow rise (Tardus)
- Sustained not displaced PMI
- Systolic thrill

$\Box S4$

Signs

Late peaking of murmur
Single S2 : Soft or absent A2
Paradoxical splitting of S2

Aortic Valve Disease

Treatment:

□ Aortic valve Replacement

Transcathter Aortic Valve Replacement Secondary Prevention of Rheumatic Fever (Prevention of Recurrent Attacks)

| Agent | Dose | Mode | | |
|---|---|------|--|--|
| Benzathine penicillin G 1 200 000 U every 4 weeks* Intramuscular | | | | |
| or | | | | |
| Penicillin V | 250 mg twice daily | Oral | | |
| or | | | | |
| Sulfadiazine 1 | 0.5 g once daily for patients 27 kg (60 lb Oral 1.0 g once daily for patients >27 kg (60 lb) | | | |
| For individuals allergic to penicillin and sulfadiazine | | | | |
| Erythromycin | 250 mg twice daily | Oral | | |
| | | | | |
| *In high-risk situations, administration every 3 weeks is justified and recommended | | | | |

Duration of Secondary Rheumatic Fever Prophylaxis

Category Duration

*

```
Rheumatic fever with carditis and<br/>residual heart disease10 y since last episode<br/>or until age 40y ,(which-<br/>(which-<br/>ever is longer), sometimes<br/>life long prophylaxis<br/>Rhumatic fever with carditis10 yrs or until age 21yrsBut no residual VHD(whichever is longer)<br/>Rheumatic fever without<br/>carditis 5 y or until age 21 y,<br/>longer)Rheumatic fever without<br/>(whichever is longer)
```