Rheumatic Mitral Valvulitis with a “Giant Vegetation”—A Case Report

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Abstract
Infective endocarditis (IE) is the infection of inner endothelial layer of the heart including the heart valves and it may present as rapidly progressive or manifest itself as subacute or chronic disease. The epidemiology of infective endocarditis has been changed over the past few decades and the incidence of IE in children in United States and Canada is 1 in 1250 pediatric hospital admissions in the early 1980s. At least 70% of infective endocarditis in children occurs with congenital heart disease whereas rheumatic heart disease in southern states of India and the degenerative mitral valve disease (myxomatous, mitral valve prolapse) in the western countries are the most underlying predisposing conditions to infective endocarditis in adolescents. The characteristic lesion of infective endocarditis is “vegetation” and a “large” vegetation >10 mm in size has been reported with an incidence of 15.9% - 62.5% in patients. The significance of vegetation size has been a subject of discussion for many years to predict the embolic episodes. Background of this case study illustrated the varying size and shape of giant vegetation attached to the anterior leaflet of mitral valve in an underlying rheumatic mitral valvulitis and its consequence of valve damage such as chordal rupture, flail leaflet and mitral regurgitation with a description of anatomic features and echocardiographic manifestations in a 10-year-old female child.

Keywords
Rheumatic Mitral Valvulitis, Infective Endocarditis, Giant Vegetation, Flail Leaflet, Mitral Regurgitation, Ping-Pong Mitral Stenosis

1. Introduction
The mitral valve apparatus consists of leaflets, subvalvular apparatus (chordae tendineae and papillary muscle), annulus and the left ventricle. It is a dynamic three-dimensional system that allows brisk left ventricular blood-inflow during diastole and ensures...
unidirectional heart pump function by sealing the left atrium from left ventricle during systole. The anterior (aortic or septal) leaflet of the mitral valve originates from the fused superior and inferior cushion tissues in the left AV (atrioventricular) canal and starts delaminating from the myocardial wall shortly thereafter. The posterior (mural) leaflet forms from the left lateral cushion. The mitral valve leaflets, chordae and papillary muscles become developed at the 15th week of gestation [1]. The anterior leaflet of the mitral valve is semicircular (trapezoid or dome-shaped) and it is larger, longer and thicker than the posterior leaflet and comprises one third of the annulus circumference and is in fibrous continuity with the left and non-coronary cusps of the aortic valve with the interleaflet triangle between the aortic cusps that abuts on to the membranous portion of the interventricular septum [2]. The posterior (mural leaflet) is quadrangular in shape and comprises two-thirds of the annular circumference. The posterior leaflet has two well defined indentations (clefts) that generally form three scallops (segments) along the elongated free edge. These leaflets are attached at their bases to the fibromuscular ring and by their free edges to the subvalvular apparatus. The commissures define a distinct area where the anterior and posterior leaflets come together at their insertion into the annulus. The leaflets are having a basal zone (connecting to the atrioventricular junction), thin central clear zone and a thick rough zone at the free edge of the leaflets. The rough zone is the main area of chordal attachment, the region of coaptation (the line of contact between the leaflets) and apposition (overlap of the leaflet free edge). The mitral valve leaflet is tri-laminar, consists of fibrosa/ventricularis, spongiosa and atrialis layers and endothelial cells cover the blood-interfering surfaces. The fibrosa is composed of dense collagen (type I—74%, type III—24%, type V—2%) [3] and it is providing the strength and stiffness to the leaflets. It is the major load bearing layer that faces the greatest pressure during valve closure, extends from the annulus into two-thirds of the leaflets and it is absent at the free edge. The spongiosa is the major component of the free edge and consists of extracellular matrix or ground substance of glycosaminoglycans and proteoglycans which are hydrophilic and attract water molecules [4]. This characteristic of water absorbent proteins causes the ground substance to expand and swell at the free edge and act as a protective buffer to ensure a tight seal along the point of apposition. Smooth muscle cells, veins and arterioles are confined to the base of the leaflets at the fibrous-myocardial junction where the leaflet inserts. Atrial myocardial cells extended into the base of the leaflets and had dense innervations and excitability, suggesting a neural regulatory mechanism [5]. The leaflets have chords (chordae tendineae), the fibrous strings that originate with highly variable branching from papillary muscle tips (heads) and insert fan-like into the ventricular aspects of the leaflets. According to the site of insertion, they are termed as primary or marginal, secondary or intermediate and tertiary or basal chords. Primary chords are inserted on the free edge of the rough zone of both leaflets and function to prevent prolapse or eversion (flail) of the leaflet margin. Secondary chords attach to the ventricular surface on the region of rough zone (i.e., body of the leaflet) and relieve tension on the valves. There is a pair of large, thick secondary chords arising from the tip of
each papillary muscle to anterior leaflet, termed as “strut” chords, which are thought to be strongest and preserving ventricular shape and function. Tertiary chords are found in posterior leaflet only and attach directly to the ventricular wall [6]. Unlike the tricuspid valve, the mitral valve does not have chords anchoring the leaflets to the ventricular septum.

Complete closure (coaptation) and correct apposition (symmetrical overlap, usually a minimum of 4 - 5 mm) of both leaflets are essential in preventing regurgitation. Leaflet coaptation depends on the balance of systolic tethering and closing forces on the valve. Tethering forces are transmitted through the LV (left ventricular) wall-papillary muscle-chordal system and keep the leaflets from protruding into the left atrium. Closing forces depend on the pressure generated by the left ventricle to close the mitral valve. Disturbances of this finely tuned spatial and temporal interplay may create an imbalance of these forces and result the valve to become regurgitant. Rheumatic valvulitis may cause mitral regurgitation by retraction of tendinous chords and leaflets as well as annular dilatation, thus compromising coaptation between the two leaflets, leading to regurgitation. Infective endocarditis can cause mitral regurgitation through chordal rupture or leaflet perforation.

**Review of Literature**

Rheumatic disease is the etiology for stenotic mitral valves in >99% of cases. The cause of pure mitral regurgitation are multiple and include floppy mitral valves, infective endocarditis, papillary muscle dysfunction, rheumatic disease, and ruptured chordae tendineae. In a study by Allen and associates [7], the three leading causes of pure mitral regurgitation were floppy (28%), rheumatic (29%), and idiopathic chordal rupture (14%). In a study by Hanson and colleagues [8], the three leading causes were floppy (70%), papillary muscle dysfunction (24%), and infective endocarditis (2%). In a study by Olson and associates [9], the reported leading causes to be floppy (38%), rheumatic (31%), and papillary muscle dysfunction (11%), but Wallen and colleagues [10] found rheumatic disease accounted for only 3% of purely regurgitant mitral valves.

Endocarditis was first described by William Osler in 1885. It is an inflammatory process that affects the endocardium and may have an infective or noninfective origin. It is uncommon in the western world (22 cases per million), but more prevalent in developing countries and a giant vegetation blocking the mitral valve orifice is uncommon and so this case had been reported.

**2. Case Report**

A 10-year-old female child was referred for echocardiographic evaluation with an apical systolic murmur. The child was having recurrent episodes of rheumatic fever (febrile illness with joint pains) at the age of 5 - 6 years and taken some treatment from the local medical practitioner, but she was not taken penicillin prophylaxis earlier. The child was remained afebrile for long period and no precipitating factors of infective endocarditis such as dental or genitourinary procedures in the past. General examination re-
revealed normal growth and development, no cyanosis and clubbing and peripheral signs of infective endocarditis such as Osler’s nodes, Janeway lesions, Roths spots and splinter haemorrhages were not present and they are relatively rare in children. Physical examination revealed a grade 3/6, blowing, high pitched, holosystolic murmur with a constant intensity and duration on dynamic auscultation and loudest at the apex with a radiation to left axilla and transmitted to the left infrascapular area and vertebral column and it is due to the flow generating the murmur is directed posterolaterally within the left atrial cavity, suggesting the murmur of mitral regurgitation due to the rupture of chordae tendineae of anterior mitral leaflet. Blood cultures were negative. Blood chemistry revealed the positive serum ASO (anti-streptolysin O) titer, suggesting a recent streptococcal infection and other parameters were normal. X-ray chest revealed moderate cardiomegaly and ECG revealed a left ventricular volume overload pattern of eccentric hypertrophy due to LV dilatation as a result of severe mitral regurgitation and a normal sinus rhythm. Transthoracic echocardiography revealed a giant vegetation “popcorn” like in Figure 1, Figure 3 and Figure 4 and “cucumber” like in Figure 2, mainly attached to base and apical portion of anterior mitral leaflet as shown in Figure 31 and manifested in various size and shapes as shown in Figures 1-37. A flail anterior leaflet with a disorganized mitral regurgitation jet as shown in Figure 21 and the posterior leaflet is embedded with vegetation and resulting in “kissing forms” as shown in Figures 13-15 in echocardiography imaging. Tricuspid valve is also thickened.

**Figure 1.** A large “popcorn” like vegetation obstructing the mitral valve—“giant vegetation” ([36], Figure 3).
Figure 2. A large “cucumber” like vegetation obstructing the mitral valve—“giant vegetation”. TV (tricuspid valve) also thickened and calcified due to rheumatic process.

Figure 3. A “popcorn” like vegetation attached to the anterior mitral leaflet moving into the left ventricle along with valve leaflet.
Figure 4. A “popcorn” like vegetation attached to the anterior mitral leaflet moving closer to the mitral valve orifice.

Figure 5. Flail anterior mitral leaflet with a pedunculated, partially calcified, vegetation mass into the left atrium (LA) ([33], Figure 13.1).
Figure 6. Healed mass of vegetation (calcified) attached with AML (anterior mitral leaflet) into LA (left atrium).

Figure 7. Vegetation seen as “rod-shaped” into LA (left atrium).
Figure 8. Vegetation seen as attached to the base of AML (anterior mitral leaflet).

Figure 9. Vegetation seen as “bileaflet structure” attached to AML.
**Figure 10.** Color Doppler imaging showing the MR (mitral regurgitation) jet as bileaflet appearance—simultaneously into LA and LV.

**Figure 11.** Vegetation seen as “basket” shaped in mitral valve orifice ([33], Figure 13.3B).
Figure 12. Vegetation seen as “ring” attached to AML (anterior mitral leaflet).

Figure 13. Vegetation seen as “kissing” type in parasternal long axis view—both AML and PML having the vegetation masses.
**Figure 14.** Vegetation seen as “kissing” type in apical four chamber view.

**Figure 15.** Vegetation seen as “kissing” type in short axis view.
Figure 16. Vegetation embedded on AML in parasternal long axis view ([33], Figure 13.7A).

Figure 17. Vegetation attached to AML (anterior mitral leaflet) showing the size in parasternal long axis view.

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Figure 18. Vegetation seen as "baby in hand" appearance on AML (anterior mitral leaflet).

Figure 19. M-mode LV function study.
**Figure 20.** M-mode LV at mitral valve level—showing vegetation on AML.

**Figure 21.** Disorganized MR jet (one component behind AML (double arrow) and the second component immediately posterior towards LA (triple arrow) of flail anterior mitral leaflet ([33], Figure 11.79).
Figure 22. Disorganized Flail MR jet—one component behind AML (anterior mitral leaflet)—highly eccentric jet.

Figure 23. Disorganized MR jet—second component posterior towards LA (left atrium).
Figure 24. The “duck” shaped MR jet—posterior towards LA.

Figure 25. The “duck” shaped MR jet—hugging the LA wall suggesting severe regurgitation.
Figure 26. Apical four chamber view showing the chaotic MR (mitral regurgitation) jet, TR (tricuspid regurgitation) jet.

Figure 27. CW (continuous wave) Doppler showing the “tiger stripe” appearance (arrows) on spectral Doppler of MR jet due to oscillating flail segments in the regurgitant stream ([33], Figure 11.85).
**Figure 28.** CW (continuous wave) Doppler showing the TR (tricuspid regurgitation) jet velocity.

**Figure 29.** Showing the functional MS (mitral stenosis—“Ping-Pong” mitral stenosis) due to the mass effect of vegetation.
Figure 30. Vegetation mass attached to AML (anterior mitral leaflet)—mitral valve in a fully opened position ([33], Figure 13.7A).

Figure 31. Vegetation mass is attached to the base and apical portion of the anterior mitral leaflet.
Figure 32. Pulsed Doppler—showing the hepatic vein systolic flow reversal suggesting severe tricuspid regurgitation.

Figure 33. Vegetation mass appearing as “bunch of plantains”—plantain tree appearance.
Figure 34. Short axis view showing the vegetation masses in the tips of mitral valve leaflets. P₁ (anterior), P₂ (middle), P₃ (posterior)—are the scallops (clefts or indentations) of PML (posterior mitral leaflet), A₁, A₂, A₃—are the corresponding segments of AML (anterior mitral leaflet). A₁ is Flail. Pc (posterior or posteromedial commissure), Ac (anterior or anterolateral commissure).

Figure 35. Short axis view showing the kissing form of vegetation masses in the LA (left atrium).
Figure 36. Subcostal view showing the vegetation on the base and tip of AML.

Figure 37. MR (mitral regurgitation) jet originating from the tip of AML (anterior mitral leaflet) at the site of attachment of vegetation into LV (left ventricle).
and calcified as shown in Figure 2 in addition to thickened and calcified mitral leaflets, suggesting an underlying rheumatic valvulitis predisposing to the formation of vegetation. The child was given 1.2 million units of intramuscular benzathine penicillin G injection as a therapeutic and initial prophylaxis dose for rheumatic fever and advised every 3 weeks for life long. Small doses of digoxin and diuretics are also prescribed and advised early surgery (mitral valve replacement).

3. Discussion

3.1. Etiopathogenesis

The mitral valve apparatus is a very complex structure and all of its components must work together for proper valve function. Mitral regurgitation can occur when any one of these elements fails through different mechanisms that affect leaflet coaptation. Intrinsincal valvular involvements by degenerative, rheumatic and infective endocarditis produce organic (primary or structural) mitral regurgitation whereas abnormal function of normal leaflets due to impaired ventricular function caused by ischemic heart disease and dilated cardiomyopathy [11], produce functional (secondary) mitral regurgitation as a result of an imbalance between tethering forces (annular dilatation, LV (left ventricular) dilatation, papillary muscle displacement, LV sphericity) and closing forces (reduced LV contractility, global LV and papillary muscle dyssynchronies and altered mitral systolic annular contraction). Papillary muscle rupture secondary to myocardial infarction defined an organic ischemic MR (mitral regurgitation). A descriptive classification of mechanism was developed by Carpentier in 1983 based upon the movement of leaflets as normal, excessive or restricted [12] as shown in Table 1 by understanding the papillary muscle-chordal-leaflet scallop relationship which has been defined anatomically by Lam et al. and it is modified by Carpentier himself in 1995 to include a mechanism of leaflet restriction during systole causing regurgitation as Type IIIb.

Kumar, et al. [13] proposed a classification (Duran classification) based on chordal insertion from the two groups of papillary muscles as lateral half of both leaflets is designated with 1 and the medial half with 2. Thus, anterior leaflet is divided into A1 with chordal crossing from the anterolateral papillary muscle and A2 with chordal crossing from the anterolateral papillary muscle and A2 with chordal crossing from the posteromedial papillary muscle. The posterior leaflet scallops are designated

Table 1. Mechanism of MR (mitral regurgitation) based on leaflet motion—Carpentier’s functional classification.

| Types    | Leaflet motion         | Mechanism                                |
|----------|------------------------|------------------------------------------|
| Type I   | Normal motion          | Annular dilation, clefts, leaflet perforation |
| Type II  | Excessive motion       | Prolapse, flail chords, ruptured papillary muscle |
| Type III | Restricted motion      |                                          |
| Type IIIa| Restriction during diastole and systole | Commissural fusion due to rheumatic heart disease |
| Type IIIb| Restriction during systole only | Chordal tethering, papillary muscle displacement |
as P₁ (anterolateral), P₂ (posteromedial) and a large middle scallop as PM. The PM is a larger scallop and subdivided as PM₁ and PM₂ based on chordal origin.

When a part of the mitral valve body protrudes into the left atrium beyond the mitral annulus and producing mild regurgitation with a preservation of coaptation is termed as “billowing valve”. It is stated that the mitral valve billows slightly into the left atrium normally and an exaggerated finding should be termed as “billowing mitral valve”, extreme form of billowing is “floppy valve” and when chordal rupture, the prolapsed mitral valve is “flail”. A morphologic abnormality with thickened leaflet (diastolic thickness > 5 mm) due to redundant tissue is termed “floppy valve”. When the coaptation line is beyond the annular plane, it is termed as mitral valve prolapse, the leaflet tip is directed towards LV and its most common phenotype is diffuse myxomatous degeneration (Barlow’s disease) in which the middle spongiosa component of the leaflet is unusually prominent and the quantity of acid mucopolysaccharide is increased. When the free edge of the leaflet is completely reversed into the left atrium (LA), the leaflet tip directed towards LA, it is termed as “flail leaflet”, usually as a consequence of chordal rupture [13]. In moderate prolapse, when the leaflet tip remains in the left ventricle, it is called as "billowing valve" and in severe prolapse when the leaflet tips bulges into the left atrium, it is called as “flail leaflet”.

Rheumatic valvulitis is characterized by variable thickening of the leaflets at their free edges, chordal fibrosis, rigidity and reduced motion of posterior leaflet in diastole and in some patients, the posterior leaflet remains in a semi-open position throughout the cardiac cycle and the friction of anterior leaflet in systole produce a false aspect of prolapse. This friction causes endothelial damage and results in platelet-fibrin deposition which is more susceptible to colonization by microorganisms. The initially sterile platelet-fibrin nidus called as nonbacterial thrombotic endocarditis (NBTE) become secondarily infected by microorganisms circulating in the blood, either from distant source of focal infection or as a result of transient bacteremia from a mucosal or skin source [14] and establishing an infectious nidus or vegetation on the endocardium and most commonly they are found at the valve closure-contact (coaptation) line on the atrial surface of the mitral valve [15]. Following successful medical therapy, the vegetative lesions may heal by a process of endothelialization of the surface phagocytes of bacterial debris, sometimes with calcification and subsequent organization by fibroblasts. Vegetation can be attached to any part of the valve and move with the leaflet but in a more chaotic (oscillating) manner and prolapse through the valve when it opens. Vegetation can prevents leaflet coaptation, and valvular retraction during the healing phase of endocarditis and result in mitral regurgitation. Large vegetation particularly at the mitral valve may result in functional valve stenosis and hemodynamic deterioration [16]. When the infection extends beyond the valve leaflet, distortion of leaflet and chordal rupture may occur, leading to severe regurgitation [17]. Ruptured mitral chordae tendineae (RMCT) are increasingly reported as an important cause of mitral regurgitation [18]. Anterior chordae tendineae rupture of the mitral valve was common in chronic rheumatic valvulitis (CRV) and infective endocarditis. In myxomatous dege-
The posterior leaflet chordal rupture is more common and the posterior leaflet becomes “flail” with a “saloon door effect”.

Rheumatic mitral regurgitation has a benign course as stated by Bland and James [19] in the analysis of the cases of rheumatic valvular lesions over a 20-year period. Levine and Friedberg have suggested that there are two groups of patients with mitral regurgitation, one group showing rapidly progressive and the other group showing a prolonged, stable and benign course. In most patients with severe primary MR (mitral regurgitation), left ventricular compensation is maintained for years, but ultimately the prolonged hemodynamic overload leads to myocardial decompensation.

**3.2. Echocardiographic Features**

Infective endocarditis is the microbial infection of the endocardial (endothelial) surface of the heart and the formation of vegetation (a variably sized amorphous mass of platelets and fibrin in which abundant microorganisms and scant inflammatory cells are enmeshed) is the hallmark of the disease and the vast majority of vegetations, however, occur on valve leaflets. The classic approach to the diagnosis of endocarditis by von Reyn does not include echocardiographic findings and it is mainly focused on pathological, clinical and laboratory evidence (positive blood cultures). In 1994, the Duke criteria for the diagnosis of infective endocarditis [20] defined “positive echocardiogram” as vegetation, an oscillating intracardiac mass, a new valvular regurgitation and microbiological evidence was established with a high sensitivity and specificity (generally about 80%). Steckelberg and Wilson [21] suggest that the incidence of endocarditis in the general population is approximately 5 cases per 100,000 persons-years. Valve redundancy and thickened leaflets (>5 mm) by echocardiography identify a population at increased risk for infective endocarditis. The relative risk of infective endocarditis in mitral valve prolapse ranges from 3.5% to 8.2%. Rheumatic heart disease is the predisposing cardiac lesion in 20% to 25% of cases in 1970’s and 1980’s [22], in Europe, it is 7% - 18% [23]. In children, it is less than one in ten cases of infective endocarditis [24] and the mitral valve is most frequently affected in women while the aortic valve in men.

Echocardiography is the only noninvasive method available for direct visualization of endocarditis-induced lesions. Echocardiographic finding in patients with infective endocarditis was initially observed by Dillon [25] and Spangler, et al. [26]. The vegetation will grow in size, either as a sessile clump or a highly mobile and even pedunculated mass with the potential for embolization. Vegetation can be detected when the valve attached mass reaches a diameter of ≥2 to 3 mm [27]. In both children and adults, 2-D echocardiography is usually the more sensitive technique with sensitivity in children up to 80% [28]. Valvular dysfunction due to tissue disruption or large obstructing vegetation can be visualized and quantitated by echocardiogram with Doppler [29]. The detection of a large eccentric jet adhering, swirling, and reaching the posterior wall of the LA is in favour of significant MR (mitral regurgitation) as shown in Figure 22, Figure 24 and Figure 25.
3.2.1. Vegetation

The most common and direct evidence of infective endocarditis is the vegetation and it begins as a microscopic focus of infection and gradually grows into a conspicuous mass. It is typically an irregularly shaped, highly mobile, echogenic mass attached to the free edge of a valve leaflet (most commonly at the coaptation line) and tends to develop on the “upstream” side of the valve leaflets (i.e., the ventricular side of aortic valve and the atrial side of mitral and tricuspid valves). They may be sessile or pedunculated, but usually has an oscillating or fluttering motion, a typical feature of most vegetations. Vegetation move with the leaflet in a more chaotic (“oscillating”) manner and it may prolapse through the valve into the LV (left ventricle) as it opens as shown in Figure 3, Figure 4 and Figure 16 and into LA (left atrium) as it closing (Figure 5 and Figure 6).

The mass of vegetation is typically homogeneous with echogenicity similar to that of the myocardium. The infectious process often alter the valvular structure and function. Extensive involvement of the leaflet may result in chordal rupture, leading to severe regurgitation as shown in Figure 21. Direct and typical signs of RMCT (ruptured mitral chordae tendineae) were chain-flail or whiplash-like changes and had an incidence of 86.7%, causing severe regurgitation and mitral chordal rupture is the leading cause of flail mitral leaflet [30]. A large vegetation may obstruct the valve orifice as shown in Figure 1 and Figure 2, sometimes termed as “obstructive-type bacterial endocarditis” and producing a functional valve stenosis (Ping-Pong mitral stenosis [31]) similar to left atrial myxoma as shown in Figure 29.

The shape and size of vegetation are quite variable and mostly it is polypoid [32]. The typical vegetation is a “sessile” or “pedunculating” valve-attached mass. A mobile vegetation showed a pedunculating part prolapsing into the ventricle as shown in Figure 3 and Figure 4 or atrium as shown in Figure 5 ([33], Figure 13.1) in a 10-year-old female child. A vegetation was considered as “definite” when shaggy echoes in the M-mode study as shown in Figure 20 and a corresponding mass without restricted valve motion in the two-dimensional echocardiogram were found as shown in Figure 16 and Figure 30 ([33], Figure 13.7). The vegetation vary in size, often being just a few mm and sometimes reaching to 2 - 3 cm. A vegetation must be at least 3 to 6 mm in size to be reliably seen. The mean size of vegetation was 0.6 mm (range 3 to 28) and vegetation >10 mm in diameter was defined as “large” and those ≤10 mm in diameter was defined as “small” and ≥15 mm is “very large”. Vegetations resulting from fungal infections (candida, aspergillus) are usually much bigger than bacterial vegetations and can be so big to be mistaken for a cardiac tumor. The large vegetations are at increased risk for embolic complications [34], especially on the anterior leaflet of the mitral valve with mobility [35]. A vegetation size of 3.2 × 4.4 cm is called as “giant vegetation” on the mitral valve with a fibrillary appearance of the mass ([36], Figure 3) as shown in Figure 1 is an important predictor of embolic phenomena in patients with infective endocarditis, causing severe mitral regurgitation as “Duck” shaped jets (Figure 24 and Figure 25), disorganized (Figure 21) and sometimes the regurgitant jet splits into two components as one into LA and the second one into LV simultaneously as bileaflet jets (Figure 10) similar to bileaflet structure of AML with vegetation masses (Figure 9) and it originates...
from tip of AML (anterior mitral leaflet) at the site of attachment of vegetation mass as shown in Figure 37. The size of the largest vegetation reported on the mitral valve in the literature in patients with bacterial endocarditis is 7 × 4 cm in a 29-year-old female at postmortem examination [37]. In a study of Nunes, et al. [38], vegetation size > 13 mm was the only independent predictor of mortality, but some studies [39] [40] did not have an increased embolic risk in patients with vegetation focused only on its presence and size and not on their location. Embolic complications may occur in infective endocarditis (20.6%) and were not more prevalent in the groups with large vegetations [41]. However, Wong, et al. [42] found an increased need for surgery in patients with a large vegetation (>10 mm).

The size and shape of vegetation vary due to curling of vegetation. The size of vegetation in this child is 35.6 × 9.3 mm as in Figure 17, 20 × 23.7 mm as in Figure 1, 32.9 × 13.9 mm as in Figure 2.

The shape of vegetation varies in this child as “popcorn” like (Figure 1, Figure 3 and Figure 4), rod-shaped (Figure 7), basket shaped (Figure 11) ([33], Figure 13.3), “baby in hand” appearance (Figure 18), “cucumber” shaped (Figure 2) and a “bunch of plantain” appearance (Figure 33), ring shaped (Figure 12), bileaflet structure (Figure 9) with bileaflet MR jet as shown in Figure 10, and kissing forms (Figure 13, parasternal long axis view, Figure 14, apical four chamber view, Figure 35, tilted apical four chamber view and Figure 15, Figure 34, short axis views).

3.2.2. “Flail” Mitral Regurgitation (MR)

The anatomic disruption of a portion of the mitral valve apparatus due to the underlying rheumatic valvulitis with predisposing infective endocarditis which form a vegetation, resulting an eccentric regurgitation jet with orientation opposite in direction of the leaflet having the anatomic defect such as “flail”. In the presence of “flail leaflet”, the mitral regurgitant spectral signal may have an atypical appearance and the flail portion oscillates in the spectral signal of regurgitant flow stream to produce a “tiger stripe” appearance as shown in Figure 27, associated with “whistling” sound on auscultation ([33], Figure 11.85). The mitral regurgitation (flail MR) jet is chaotic as shown in Figure 26, highly eccentric (Figure 22) and disorganized with one component behind the anterior mitral leaflet and the second component directed posterior immediately as in Figure 21 ([33], Figure 11.79).

The severity of eccentric MR is underestimated because of coanda effect. If the regurgitant jet area fills <20% of the left atrium, it is mild and >40% indicates severe regurgitation. The vena contracta (the neck or narrowest portion of the jet), typically imaged perpendicular to the commissural line in parasternal long axis and apical four chamber views is well defined in both central and eccentric jets, but not in chaotic, disorganized jets due to flail leaflets. Its width < 3 mm indicates mild MR, >7 mm defines severe MR and a mean value of >8 mm indicates severe functional MR. The flow convergence method based on PISA (proximal isovelocity surface area) may not applicant for eccentric and multiple jets or complex and elliptical regurgitant orifices to assess the
severity of mitral regurgitation. The adaptation of LV to the increased volume overload is reflected by LV dimensions and ejection fraction. In chronic compensated phase, the forward stroke volume is maintained through an increase in LV ejection fraction >65% and the patient could be asymptomatic. In chronic decompensated phase of MR, the forward stroke volume decreases and the LA pressure increase significantly. The patient may be still asymptomatic and the LV ejection fraction may be in the low normal range despite the presence of significant muscle dysfunction. The contractile function decreases silently and become irreversible. In the current guidelines, surgery is recommended in asymptomatic patients with severe organic MR when the LV ejection fraction is ≤60%. However, in acute stage, the LV ejection fraction increases in response to the increased preload. The end-systolic diameter is less preload dependent than the ejection fraction and it may be more appropriate to monitor the global LV function. The end-systolic diameter > 45 mm also indicate the need for mitral valve surgery [43]. In this child, the LVESD (end-systolic diameter) is 30.7 mm and the ejection fraction (EF) is 66% as shown in Figure 19. New parameters are currently available for a better assessment of LV function. A systolic tissue Doppler velocity measured at the lateral annulus < 10.5 cm/s has been shown to identify significant LV dysfunction and to predict post-operative LV function in patients with asymptomatic organic MR [44]. Strain imaging allows a more accurate estimate of myocardial contractility than tissue Doppler velocities. In MR, strain has been shown to decrease even before LV end-systolic diameter exceeds 45 mm [45]. A resting longitudinal strain rate < 1.07/s (average 12 basal and mid segments) is associated with subclinical latent LV dysfunction [46] and a global longitudinal strain <18.1% has been associated with postoperative LV dysfunction [47]. The left atrium dilates in response to chronic volume and pressure overloads. LA remodelling (diameter > 40 - 50 mm) may predict the onset of atrial fibrillation and poor prognosis in patients with organic MR [48]. The excess regurgitant blood entering in the LA may induce acutely or chronically a progressive rise in pulmonary pressure and the presence of TR (tricuspid regurgitation) as shown in Figure 26 and Figure 28 permits the estimate of systolic pulmonary arterial pressure and mitral valve surgery is recommended when it is >50 mmHg at rest and LA reverse remodelling may occur after surgery. The severe TR may cause a decrease in hepatic vein systolic velocity and systolic flow reversal may occur as shown in Figure 32 and its sensitivity is 80% [49]. The TR (tricuspid regurgitation) jet velocity in this child is 4.03 m/s as shown in Figure 28 which corresponds to a systolic pulmonary artery pressure of 65 mmHg.

3.3. Treatment

3.3.1. Pharmacological Therapy
Pharmacological therapy aims to alleviate symptoms and to prevent the progression of LV dysfunction. Afterload reduction is of particular benefit in the management of both acute and chronic forms of mitral regurgitation [50]. Afterload reduction with sodium nitroprusside is life saving in acute conditions such as chordal or papillary muscle rupture. It stabilizes the patient by reducing the impedance to the LV ejection and thus, the
regurgitant volume and LA pressure decrease. Dobutamine may be administered along with nitroprusside if hypotension exists. Afterload reduction in chronic, severe MR with vasodilators such as angiotensin converting enzyme inhibitors are beneficial in presence of heart failure, but not beneficial in its absence [51]. In addition to diuretics, digitalis glycosides are indicated in patients having severe MR and heart failure, particularly with established atrial fibrillation along with anticoagulants. IE (infective endocarditis) prophylaxis was advised during surgical procedures in these patients.

3.3.2. Surgical Therapy

Asymptomatic patients with severe MR having excellent ventricular function (EF > 70%, ESD (end-systolic diameter) < 40 mm) can be safely followed by a “watchful waiting approach” until symptoms, LV dysfunction (EF ≤ 60%, ESD ≥ 45 mm) and pulmonary hypertension develops. Mitral valve surgery is the only treatment for MR which provides substantial relief of symptoms and prevent the development of heart failure [52]. The normal function of the mitral valve apparatus “primes” the left ventricle for normal contraction. Operative procedures may interfere the annular-chordal-papillary muscle continuity, results in postoperative LV dysfunction and so the preservation of these structures are now considered as a critical feature of MVR (mitral valve replacement) [53]. Reconstructive procedures are carried out in degenerative MR due to mitral valve prolapse and chordal rupture as well as in ischemic functional MR [54]. Percutaneous catheter-based mitral valve repair procedures such as leaflet edge-to-edge repair and mitral annular reduction are currently under clinical evaluation and the preliminary results are encouraging [55]. The severely deformed valves in rheumatic heart disease are not suitable for reconstructive surgery and they require MVR [56] and the operation should be desirable before they develop marked LV dysfunction [57] and serious symptoms since severe LV dysfunction(EF < 30%) may cause high perioperative mortality [58]. Excellent survival is observed in patients with ESD < 45 mm, EF ≥ 60% and the 5-year survival rate is 40% in ischemic MR and 75% in rheumatic MR. Intermediate outcome may occur when LV ESD is 45 - 52 mm and the ejection fraction between 50% - 60%. Poor outcome is associated with values below these limits.

Surgery may be considered in mobile vegetation > 10 mm in size since the incidence of systemic embolization are increased (33%) compared to those with smaller size (19%) and particularly those on anterior mitral leaflet are uniquely associated with embolic episodes. Urgent surgery is indicated when a large vegetation is associated with embolic events, heart failure and persistent infection [59]. In patients with valvular dysfunction and in whom the infection is controlled with antibiotic therapy and cardiac function is compensated, surgery may be delayed until antimicrobial therapy has been completed. Early surgery is appropriate in patients with large vegetation who may need valve replacement in future since larger vegetations are associated with increased risk of mortality and embolization [60] [61]. Some authors suggest that the causative microorganism is associated with embolic complications rather than the size of vegetation [62].

Since this child is having a giant vegetation with disorganized severe mitral regurgi-
tation due to flail anterior mitral leaflet as the result of chordal rupture, the child was advised MVR (mitral valve replacement) with mechanical protheses along with removal of vegetation and to continue penicillin prophylaxis with anticoagulant therapy for life long. Since the vegetation is healed, partially calcified and an organized mass, the antimicrobial regimen for infective endocarditis was not administered. The anticoagulant therapy is not preferred at this moment since the child had no embolic episodes and the role of prophylactic anticoagulant therapy is questionable in large vegetation in the absence of atrial fibrillation. The child was advised small doses of digoxin and diuretics along with penicillin prophylaxis.

3.4. Outcome

Several factors play a role in the prognosis of IE (infective endocarditis). Costa, et al. described the clinical and echocardiographic scores as the predictors of mortality in infective endocarditis. Age > 40 years (4 points), class IV heart failure (cardiogenic shock) (4 points), uncontrolled sepsis (6 points), conduction disease (5 points), arrhythmias (8 points), a valve with excessive damage (5 points), large and mobile vegetation (4 points) [63]. Mortality rates for scores below 10 were 5.26% and the scores over 20 were 78.9%. The child is having a large and mobile vegetation (4 points), extensive valve damage (5 points) and according to this study, the mortality rate of this child is 5.26%.

3.5. Screening of Population

Transthoracic echocardiographic screening revealed a “popcorn” like active mobile vegetation attached to the flail anterior mitral leaflet and producing chordal rupture with severe, disorganized mitral regurgitation as shown in Figures 38-43 in a 5-year-old febrile (one week duration) boy having an underlying rheumatic mitral valvulitis with a grade 3/6 holosystolic murmur in the apex (radiating to axilla and back) and had an intermittent diastolic sound (“vegetation flop”) coinciding with the physical movement of vegetation mass across the mitral valve orifice [64] at the apex. The vegetation is attached to the base of AML since the basal portion has some vascular supply at the fibrous-myocardial junction where the leaflet inserts and so the basal region is more susceptible to vegetation formation from the blood-borne infections. A small “popcorn” like mobile vegetation attached to the junction of anterior mitral leaflet and chordae is detected by echocardiography in a 46-year old febrile female as shown in Figure 44 due to friction and trauma.

A “sessile” vegetation attached to the atrial side of posterior mitral leaflet, producing a “flail” PML with a “saloon door” effect and a “bileaflet” severe mitral regurgitation into the left atrium in a 95-year-old asymptomatic female was shown in Figures 45-48. Characteristically, the bileaflet flail PML (posterior mitral leaflet) MR (mitral regurgitation) jet is directed into left atrium only as in Figure 46 and Figure 48 whereas the bileaflet flail anterior mitral leaflet MR jet is directed into both LA and LV simultaneously as shown in Figure 10. A “sessile” vegetation had to be completely attached to the valve as shown in Figure 49 and Figure 50 in a 63-year-old male, in which a large vegetation...
is attached to the atrial side of anterior mitral leaflet ([33], Figure 13.3-A), producing mitral regurgitation as shown in **Figure 51**. A “popcorn” like “giant” vegetation, attached to the anterior mitral leaflet which prolapse with mild mitral regurgitation in a 49-year-old febrile male was illustrated in **Figures 52-62**.

**Figure 38.** A “popcorn” like vegetation attached to AML in a 5-year-old febrile boy—moving into the LV along with the leaflet. PML (posterior mitral leaflet) is also having the vegetation.

**Figure 39.** Vegetation masses seen in both AML and PML in the 5-year-old febrile boy—active vegetations.
**Figure 40.** Chordal rupture and “flail” AML in the 5-year-old boy.

**Figure 41.** Vegetation mass attached to the flail AML—prolapsing into LA.
Figure 42. Vegetation mass attached to the base of AML in the 5-year-old boy.

Figure 43. Chordal rupture producing severe disorganized MR jet as one component behind AML and the second component immediately posterior due to flail AML in the 5-year-old boy.
Figure 44. A “popcorn” like small vegetation seen at the junction of AML and chordal attachment in a 46-year-old febrile female.

Figure 45. A sessile vegetation attached to PML (posterior mitral leaflet) in a 95-year-old asymptomatic female—parasternal long axis view.
Figure 46. A sessile vegetation in PML-producing "bileaflet MR" in a 95-year-old asymptomatic female (mitral regurgitation)—parasternal long axis view.

Figure 47. A sessile vegetation attached to PML—leading to chordal rupture and flail PML with "saloon door effect" in a 95-year-old asymptomatic female—apical four chamber view.
Figure 48. A sessile vegetation attached to PML-producing “bileaflet MR” in apical four chamber view in a 95-year-old asymptomatic female.

Figure 49. “Sessile” vegetation attached to AML in a 63-year-old male ([33], Figure 13.3A).
Figure 50. "Sessile" vegetation showing calcification in a 63-year-old male.

Figure 51. "Sessile" vegetation causing mitral regurgitation in a 63-year-old male.
Figure 52. A “popcorn” like “giant” vegetation attached to anterior mitral leaflet (AML) in a 49-year-old febrile male.

Figure 53. A “popcorn” like vegetation—seen as an active giant vegetation at the tip of AML in a 49-year-old febrile male.
Figure 54. A “popcorn” like vegetation—seen as a calcified mass at the tip of AML in a 49-year-old febrile male.

Figure 55. A “popcorn” like vegetation-prolapsing into LV along with the anterior mitral leaflet in a 49-year-old febrile male.
Figure 56. A “popcorn” like vegetation mass attached to the tip of AML in a 49-year-old febrile male.
Figure 57. (a) A “popcorn” like vegetation mass into the LV cavity in 49-year-old febrile male; (b) A “popcorn” like vegetation mass in AML at mitral orifice in a 49-year-old febrile male.

Figure 58. The vegetation is attached to AML-apical four chamber view in a 49-year-old febrile male.
**Figure 59.** The vegetation mass, attached to AML (anterior mitral leaflet) seen as partially calcified mass (arrow) near the mitral orifice in a 49-year-old febrile male.

**Figure 60.** The AML (anterior mitral leaflet)—showing “prolapsed” (vertical arrow) with mild mitral regurgitation (MR) (left horizontal arrow) in a 49-year-old febrile male.
Figure 61. A "giant" vegetation attached to AML (anterior mitral leaflet) seen near the aortic orifice (LV outflow tract) in a 49-year-old febrile male.

Figure 62. A "giant" vegetation attached to AML (anterior mitral leaflet) seen as moving towards aortic orifice in a 49-year-old febrile male.
4. Conclusion

A giant “popcorn” like vegetation attached to the anterior mitral leaflet and presented with various shapes in a 10-year-old female child was described by Transthoracic 2D echocardiography imaging. The child had underlying rheumatic involvement of the mitral valve and the infective endocarditis, causing chordal rupture leading to flail anterior mitral leaflet with severe mitral regurgitation. The vegetation functionally obstructs the mitral orifice and results in “ping-pong” mitral stenosis due to mass effect. The child remained asymptomatic without any embolic episodes and heart failure symptoms on follow up of two years. The “popcorn” like vegetations attached to AML (anterior mitral leaflet) in a 5-year-old boy and in a 49-year-old male were detected. The “sessile” vegetations attached to AML in a 63-year-old male with severe mitral regurgitation and attached to PML (posterior mitral leaflet) in a 95-year-old asymptomatic female with bileaflet severe mitral regurgitation were also found by Transthoracic 2D echocardiographic imaging at this region of Thoothukudi in India.

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