Mechanisms, features, and significance of diastolic mitral regurgitation: a case series

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Background
Diastolic mitral regurgitation (DMR) is a type of functional mitral regurgitation. Its occurrence in the diastolic phase of cardiac cycle renders DMR an easily ignored entity. Confusing it with systolic mitral regurgitation occasionally happens. The reversal of left atrioventricular pressure gradient during diastole and the incomplete closure of mitral valve are the essential conditions for DMR. Diastolic mitral regurgitation develops under various situations, where the mechanisms of diastolic reversal of left atrioventricular pressure gradient differ.

Case summary
Patient 1 was a 50-year-old man diagnosed with 2:1 second-degree atrioventricular block (AVB). Patient 2 was a 70-year-old man diagnosed with first-degree AVB. Patient 3 was a 66-year-old man diagnosed with atrial fibrillation with long intermission and occasional atrial flutter with unequal conduction. Patient 4 was a 54-year-old woman diagnosed with dilated cardiomyopathy with complete left bundle branch block. Patient 5 was a 36-year-old man diagnosed with severe acute aortic regurgitation secondary to subacute bacterial endocarditis.

Discussion
Although the degree of DMR is relatively mild, its appearance generally prompts further clinical considerations. The appreciation of DMR has an incremental value for diagnosing and evaluating the underlying cardiovascular disease.

Keywords
Case series • Diastolic mitral regurgitation • Functional mitral regurgitation • Mechanisms

Learning points
• Diastolic mitral regurgitation is easily ignored because of the different characteristics from systolic mitral regurgitation.
• Diastolic mitral regurgitation could be caused by different mechanisms.
• It is of great significance to recognize diastolic mitral regurgitation because it may offer incremental information for clinical decision-making.

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**Introduction**

Mitral regurgitation (MR) is the most common valve insufficiency and is classified as primary or functional. Based on the phase in cardiac cycle, functional MR can be further divided into systolic mitral regurgitation (SMR) and diastolic mitral regurgitation (DMR). Diastolic mitral regurgitation is a rare phenomenon which appears with or without structural heart disease. Although DMR has been noticed for decades and plenty of case reports have been published, far less attention is paid to DMR than to SMR. Moreover, the rarity and occasional co-existence with specific heart disease of DMR, along with its differing clinical implications and haemodynamic consequences, necessitate the differentiation between DMR and SMR. This usually requires multiple echocardiography views and electrocardiogram (ECG) recordings. The recognition of DMR supplements an accurate diagnosis and evaluation of cardiovascular disease. In this article, we will discuss the mechanisms, echocardiographic features, and clinical significance of DMR by five cases.

**Timeline**

Baseline characteristics of the five patients included in this case series.

| Patient | Age | Gender | Diagnosis | Relevant mechanism |
|---------|-----|--------|-----------|--------------------|
| 1       | 50  | Male   | 2:1 second-degree atrioventricular block (AVB) | ‘Overdue’ left ventricle (LV) systole |
| 2       | 70  | Male   | First-degree AVB | ‘Overdue’ LV systole |
| 3       | 66  | Male   | Atrial fibrillation with long intermission and atrial flutter | ‘Overdue’ LV systole |
| 4       | 54  | Female | Dilated cardiomyopathy with complete left bundle branch block | LV systolic asynchrony |
| 5       | 36  | Male   | Severe acute aortic regurgitation (subacute bacterial endocarditis) | Elevation of LV diastolic pressure |

**Case presentations**

**Patient 1**

A 50-year-old man presented to our hospital complaining of exertional dyspnoea. Laboratory test showed his N-terminal pro-brain natriuretic peptide (NT-proBNP) was within normal range. Electrocardiogram showed 2:1 second-degree atrioventricular block (AVB). Transthoracic echocardiography (TTE) showed no evidence of pulmonary hypertension and an enlarged left atrium (LA) and DMR. The characteristics of DMR are shown in Figure 1 and Video 1. A pacemaker was implanted and atrioventricular rhythm was normalized thereafter. Diastolic mitral regurgitation disappeared immediately. During a 2-year follow-up, the pacemaker functioned well, and there was no DMR anymore.

**Patient 2**

A 70-year-old man was diagnosed with first-degree AVB in his routine medical examination. His medical history was unremarkable for cardiovascular disease and he had no obvious clinical symptoms. Laboratory test showed his NT-proBNP was within normal range. Transthoracic echocardiography showed mild SMR and DMR. The characteristics of DMR are shown in Figure 2. There was no clinical intervention for the patient. During a 1-year follow-up, first-degree AVB and the DMR persisted.

**Patient 3**

A 66-year-old man presented to our hospital complaining of recurrent palpitation over 3 months. He had a history of hypertension for 8 years and diabetes mellitus for 6 years. The laboratory test revealed a slightly elevated level of NT-proBNP at 327.6 pg/mL (ref. range: 0–100 pg/mL). Electrocardiogram revealed atrial fibrillation with long intermission and occasional atrial flutter with unequal conduction. Transthoracic echocardiography showed atrial enlargement, DMR, and mild SMR. The characteristics of DMR are shown in Figure 3. This patient received medication for anticoagulation and control of ventricular rate. In the recent follow-up, the arrhythmia persisted, so did DMR.

**Patient 4**

A 54-year-old woman presented with 1-year history of exertional dyspnoea. The laboratory test revealed an elevated level of NT-proBNP at 1828.5 pg/mL. Electrocardiogram showed complete left bundle branch block (CLBBB). Transthoracic echocardiography demonstrated a dilated left ventricular (LV) with global hypokinesis and asynchrony. His left ventricular ejection fraction (LVEF) was 32%. Subsequent coronary computed tomography angiography revealed normal coronary arteries. A diagnosis of dilated cardiomyopathy (DCM) was made. Colour flow imaging revealed DMR and moderate SMR. The characteristics of DMR are shown in Figure 4 and Video 2. Medical therapy with betablockers, angiotensin-converting-enzyme inhibitors, and diuretics was initiated. The patient did not receive device-based therapy. His LVEF had improved to 40% after 1 and a half year of medication but DMR remained.

**Patient 5**

A 36-year-old man was admitted to our hospital for shortness of breath and lower extremity oedema for the previous 2 weeks. This patient reported an history of intermittent fever for the previous 2 months. On admission, his heart rate was 103 b.p.m., blood pressure was 122/47 mmHg, and temperature was 37.2 °C. Cardiac examination revealed a pandiastolic murmur in the aortic area. Labs showed an elevated level of NT-proBNP at 4492.0 pg/mL. Transthoracic echocardiography demonstrated bicuspid aortic valve...
and severe aortic regurgitation (AR) secondary to vegetations and prolapse of aortic valve. In addition, there was DMR. Subsequent blood culture grew Streptococcus gordonii and confirmed the diagnosis of subacute bacterial endocarditis. Emergency aortic valve replacement was performed on the 4th day for aggravated heart failure.

Transoesophageal echocardiography findings were similar to TTE. The characteristics of DMR are shown in Figure 5 and Video 3. After aortic valve replacement, DMR disappeared immediately. The most recent TTE showed the prosthetic valve functioned well, and there was no DMR.
Discussion

Essential conditions for diastolic mitral regurgitation

Blood flow is driven by pressure gradient, moving from higher pressure area to lower pressure area. Under normal conditions, a negative pressure gradient from LV to LA during diastole keeps a forward blood flow through mitral valve (MV), which then fills the LV. A reversal of this pressure gradient establishes the haemodynamic foundation for DMR, which can be achieved or maintained during various stages of diastole through different mechanisms.2,3

Full closure of the normal MV requires effective LV contraction with enough closing force, coordinated papillary muscles...
mechanical activation and normal mitral annular sphincteric contraction. Normally, after LA contraction, LV diastolic pressure will increase above LA pressure and drive MV leaflets to approximate each other. However, this small pressure gradient is not enough to achieve a full closure in the lack of effective LV contraction. Hence, the MV will keep an incomplete closure state during diastole and thus provide a potential passage for DMR. In addition, SMR patients are predisposed to DMR in the same manner.

Therefore, the reversal of left atrioventricular pressure gradient and the incomplete closure of MV are the essential conditions for DMR. Diastolic mitral regurgitation may be caused by a variety of diseases, and the mechanisms encompass ‘overdue’ LV systole, LV systolic asynchrony, and increased LV diastolic pressure. These mechanisms can induce DMR separately, or work in concert to aggravate the degree of DMR.
Overdue left ventricular systole
A delayed LV contraction mostly coincides with a certain kind of arrhythmia, such as AVB or rapid atrial arrhythmia with long intermission. When these arrhythmias conform to certain conditions, DMR will be induced based on the two aforementioned essential factors. It is worth mentioning that the ‘overdue’ LV systole is the most common mechanism of DMR.

Atrioventricular block
Diastolic mitral regurgitation may be observed during any degree of AVB where LA contraction is not followed by LV contraction immediately. Atrioventricular block can be roughly summarized as conduction failure or conduction prolongation of the sinus P-wave. In these two conditions, the mechanisms of DMR are similar.

The most classic example of sinus P-wave conduction failure-induced DMR is 2:1 second-degree AVB. After a conducted P-wave, LV contraction is followed by its active relaxation and an increase in LV pressure, while the next blocked P-wave causes LA contraction to further elevate the LV diastolic pressure. Subsequent LA relaxation then coincides with the elevated LV diastolic pressure. The atrioventricular pressure gradient will reverse once the LA pressure falls below that of the LV. Meanwhile, a properly timed LV contraction does not kick in, which leaves the MV leaflets incompletely closed. As a consequence, during the extended diastole of LV, DMR appears immediately after LA contraction.

In situation of sinus P-wave conduction prolongation, such as first-degree AVB, prolonged P-R interval delays LV contraction. Similar to 2:1 second-degree AVB, DMR may appear in the extended LV diastolic phase between LA contraction and LV contraction.

For other AVB, namely the rest second-degree and third-degree AVB, similar mechanism plays a role in the formation of DMR. Like SMR, DMR also increases the volume load of LA, while the degree is relatively slight because of the low-pressure gradient. However, inability to distinguish DMR from SMR and sometimes a confusingly large jet area of DMR can together lead to erroneous judgement of the severity of regurgitation. Diastolic mitral regurgitation caused by AVB is, in general, a benign phenomenon which can mostly be improved after treatment of AVB.

Atrial arrhythmia
Rapid atrial arrhythmias (e.g. atrial flutter and atrial fibrillation) with long intermission are also common causes of DMR, both of which have extremely rapid atrial frequency. Once a relatively long RR interval presents, LA contractions can repeat several times between the two successive LV contractions. After every LA contraction, as in AVB, LV diastolic pressure increases while LA relaxes shortly after. Thus, one or more episodes of left atrioventricular pressure gradient reversal may happen in diastole and result in several separate occurrences of DMR until next LV contraction.

In another special situation, blocked atrial premature beat may induce DMR under certain conditions, where the electrocardiographic findings around this cardiac cycle is very similar to 2:1 second-degree AVB. A blocked atrial premature beat appears following the previous LV contraction and the atrioventricular pressure gradient may reverse immediately after this LA contraction, which would generate a one-time DMR.

In these atrial arrhythmias, DMR is also a benign phenomenon because of its low-pressure gradient and short duration and can be mostly relieved after the treatment of atrial arrhythmia.

Left ventricular systolic asynchrony
Diastolic mitral regurgitation associated with LV systolic asynchrony is another interesting mechanism, and this pathological state is relatively common in DCM and CLBBB.

Left ventricular systolic dyssynchrony in DCM is due to inter- or intra-ventricular conduction delay, which can lead to asynchronous contraction of different LV wall regions. A few myocardial segments may perform post-systolic shortening after the closure of aortic valve, which might cause paradoxical increase of LV pressure during early diastole. The subsequent elevated pressure of LV might lead to an inversion of pressure gradient from LV to LA in early diastolic phase. Moreover, this pressure gradient, far below that of previous overall LV contraction process, may not be enough to drive MV towards full closure. Besides, in certain DCM, malfunctioning papillary muscles due to uncoordinated activation of corresponding segments

Video 2 Dilated cardiomyopathy with DMR.

Video 3 Severe aortic regurgitation with DMR.
leads to geometrical changes of MV and increased tethering. This process may occur in both LV overall contraction and post-systolic contraction situations. Taken together, DMR would occur in DCM patients during early diastolic phase.

This phenomenon may also be observed in CLBBB under a similar mechanism, especially in DCM combining with CLBBB.

It is worth noting that the pressure gradient reversal caused by post-systolic contraction may postpone the MV opening. Such phenomenon may be the net result of the discordant movement between segments, where partial segmental relaxation coexists with partial segmental post-systolic contraction. Therefore, although this mild DMR cannot increase the volume load of LA markedly, the early diastolic filling process of LV is hindered, and LV filling time in whole cardiac cycle is shortened to a large extent. Hence, DMR in early diastole indicates prominent LV systolic asynchrony and limited early diastolic filling. Cardiac resynchronization therapy (CRT), which helps to re-establish the synchronous contraction of different LV wall regions, may also improve the effective LV diastolic filling.6,13

Elevation of left ventricular diastolic pressure

The increase in LV diastolic pressure is another special cause of DMR. For instance, in patients with severe AR, especially acute AR, haemodynamic disturbance during diastole is the culprit of DMR. This phenomenon is relatively rare in patients with chronic AR due to LV remodelling as a result of volume load adaptation.15

In severe AR, LV accepts blood not only from LA but also from the aorta during diastole. Thus, the AR ‘overfills’ the LV and makes the filling pressure of LV rapidly increase and possibly exceed the LA pressure in late diastole, which may produce presystolic incomplete MV closure. As a consequence, in late diastole, DMR may appear and last for a period of time.

It is noteworthy that this reversal of pressure gradient impedes the transient opening of MV by atrial contraction, adds to LA volume overload and aggravates pulmonary oedema. This highlights the importance to recognize DMR in severe AR as it indicates the acuteness of AR and the need for urgent surgical intervention, while DMR in itself is a reversible process as it will disappear after successful aortic valve replacement.16

Other possible causes of the elevation of LV pressure in diastole may also lead to the formation of DMR, such as mild or moderate AR, LV diastolic dysfunction, or LV blood pooling in the setting of dynamic left ventricular outflow tract obstruction with apical hypokinesia.17–19 Although not sufficient to induce DMR separately, they may aggravate the degree of DMR in other situations (e.g. AVB) or combinatorially cause DMR.

Conclusions

Despite an uncommon finding, DMR can occur in a variety of diseases. Echocardiography with colour Doppler, spectral Doppler, and M-mode, combined with real-time ECG, represents the most useful imaging modality to identify DMR, because it enables visualization of the direction, velocity, and phase of MR. Understanding the various forming mechanisms of DMR under different disease state helps explain its relatively benign nature and the possible alternative therapeutic effect of CRT. Although the degree of DMR is relatively mild and poses insignificant regurgitant volume load to the heart, differentiation of DMR from SMR prevents overestimation of regurgitation and unnecessary clinical intervention and signifies urgent surgical correction when complicating severe AR.

Lead author biography

Quan Li is a graduate of Nanjing Medical University, Jiangsu Province, China in 2011. Currently, he worked at Department of Echocardiography, Zhongshan Hospital, Fudan University, Shanghai. His research interest is mainly focused on multimodal cardiac imaging, in particular, in valvular heart diseases and haemodynamics.

Supplementary material

Supplementary material is available at European Heart Journal - Case Reports online.

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Slide sets: A fully edited slide set detailing this case and suitable for local presentation is available online as Supplementary data.

Consent: The author/s confirm that written consent for submission and publication of this case report including image(s) and associated text has been obtained from the patients in line with COPE guidance.

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