Cleft-like indentations in myxomatous mitral valves by three-dimensional echocardiographic imaging

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ABSTRACT

Objectives Cleft-like indentations (CLI) are deep separations between scallops of the mitral posterior leaflet observed in myxomatous mitral valve disease (MMVD), but their diagnosis, mechanisms and implications are unknown. Using 3D transoesophageal echocardiography (3DTOC), we aimed at assessing diagnostic accuracy and defining mechanisms of CLI in patients undergoing surgery for MMVD.

Methods 3DTOC of mitral valve was acquired in 49 patients with MMVD and severe regurgitation prior to valve repair. Qualitative review compared 3DTOC diagnosis of CLI with surgical inspection. Mitral, annular and leaflet dimensions were quantified with dedicated software and compared between those with and without CLI.

Results Diagnosis of CLI was made by 3DTOC in 17 (35%) while none was identified by 2D and was confirmed in 15 (88%) by surgical inspection. Mechanistically, LV diameters and mitral regurgitant volume (RVol) were similar with and without CLI (p=0.49). Conversely, mitral annulus was smaller with CLI (anteroposterior diameter 42.2±7.1 vs 47.0±7.5 mm, p=0.04; circumference 133±16 vs 148±7.5 mm, p=0.04; area 1289±326 vs 1619±427 mm², p=0.01). Prolapso volume tended to be smaller with CLI (1.9±1.2 vs 4.0±4.3 mL, p=0.06) involving single posterior scallop at surgery (82% vs 44%, p=0.007) with smaller 3DTOC leaflet area (1574±409 vs 2019±652 mm², p=0.01). During valve repair, surgical closure of all surgically diagnosed CLI was required.

Conclusions Posterior leaflet CLI are frequent in MMVD, are identified by 3DTOC with high accuracy and require closure during valve repair. CLI are mechanically not related to excess annular enlargement or excess prolapse. Conversely, CLI occur in the context of single scallop prolapse with tissue paucity causing excess separation of scallops. These 3DTOC data enhance diagnostic and mechanistic comprehension of the diversity of MMVD phenotypical presentation.

INTRODUCTION

Myxomatous mitral valve disease (MMVD) is the leading cause of organic severe mitral regurgitation (MR) in western countries and frequently requires surgical treatment.1 Mitral valve repair is the procedure of choice for patients with MMVD because it can restore normal life expectancy when timely performed.2 However, the success of mitral valve repair, defined by the ability to perform the procedure during surgery and by long-term durability,3 depends on the congruence of surgeon’s expertise with specific lesions to be repaired.4 Predicting successful repair is crucial to indicating surgery in most recent guidelines5 and demands accurate mitral anatomic assessment and characterisation of leaflets’ lesions.6

3D transoesophageal echocardiography (3DTOC) profoundly changed the ability to visualise the complete mitral valve anatomy in vivo7,8 while gaps in mitral tissue are better diagnosed. True posterior leaflet clefts, cleaving it in two equal remnants, while long-known,9 are rare, but have recently been diagnosed in vivo with 3DTOC.10,11 Lately, another feature of MMVD was described: deep ‘clefs’ of posterior mitral leaflet,12 which do not really cleave the leaflet, but occupy the position of normal indentations between its three scallops. Normal indentations are usually not deep, but abnormal indentations, although different from true clefs,9 form when they are deep cleft-like indentations (CLI) that can be associated with MR13 and failure of mitral repair.14

These recently recognised CLI have been hypothesised as linked to mitral prolapse extent12 and not to MR per se. Whether annular enlargement, tissue excess or prolapse severity are associated with CLI presence is unknown, and little is known about CLI mechanism. Therefore, we initiated a prospective 3DTOC study of patients with MMVD undergoing mitral repair for MR, aiming at defining accuracy of CLI preoperative diagnosis and at defining 3D MMVD morphological features independently linked to CLI presence. Our working mechanistic hypothesis was that excess annular enlargement, by separating posterior mitral scallops, was CLI’s main causal determinant.

METHODS

The study included 49 consecutive patients referred for elective surgical mitral valve repair for MMVD with high-quality 3DTOC quantifiable. Patients were enrolled prospectively if they had (1) severe MR due to MMVD, (2) indicating mitral valve repair, (3) comprehensive transthoracic Doppler echocardiography performed at Mayo Clinic during the same episode of care, (4) 2D and 3D intraoperative transoesophageal echocardiography (TEE) of mitral valve and (5) 3D datasets of sufficient quality to quantify mitral annulus and leaflets alterations. Exclusion criteria were contraindications to TEE,
uncontrolled atrial fibrillation, mitral stenosis, moderate-to-severe aortic valve disease, moderate-to-severe tricuspid regurgitation or associated pericardial or congenital heart disease. Patients with MMVD associated with connective tissue disorders, Marfan syndrome, hypertrophic cardiomyopathy and rheumatic heart disease were also excluded. The study was approved by Mayo Institutional Review Board and was deemed low-risk, requiring only verbal consent.

**Doppler and 2D echocardiographic measurements**

Comprehensive Doppler and 2D echocardiography data were prospectively recorded preoperatively. LV, left atrium (LA) dimensions and LV EF were measured by 2D echocardiography. MR was quantified as effective regurgitant orifice (ERO) and regurgitant volume (RVol).

**3DTOC qualitative and quantitative image analysis**

TEE was performed intraoperatively, after initial anaesthesia induction and endotracheal intubation, before cardiopulmonary bypass or thoracic incisions. The ultrasound platform was RT3DE imaging probe (model X7-2t) and iE33 echocardiography imaging platform (Philips Medical Systems, Bothell, Washington, USA).

From mid-oesophageal position, live 3DTOC zoom of mitral valve included aortic valve and entire mitral annulus throughout the cardiac cycle. For mitral qualitative assessment, comprehensive imaging was obtained by standard 2DTEE and by 3DTOC after cropping. 3DTOC was recorded both as live 3D (single-beat) and as multi-beat full volume datasets, and imaging was optimised to enhance physical and temporal (8–12 Hz) resolution. For 3DTOC interpretation, both atrial and ventricular views of the mitral valve were analysed in detail, with swivel examination of the entire valve to examine all three scallops of posterior leaflet and of indented zones between scallops. Diagnosis of CLI by echocardiography was defined as a visible tissue defect seen during systole, occupying the interscallop region of the posterior leaflet—not plicatures of leaflets’ free edge at other location—(figure 1) and extending at least half depth of the adjacent mitral scallop. The ventricular aspect of mitral valve was also examined for careful assessment of mitral scallops with possible visualisation of CLI from this orientation (figure 2). The diagnostic views were stored for off-line interpretation before left atrial opening by the surgeon who directly examined the mitral valve (figure 1).

For quantitative analysis of mitral valve, full-volume 3D datasets were digitally stored and transferred to a workstation with Q-Laboratory (QLAB) Mitral Valve Quantification Software (Philips Medical Systems) for off-line analysis by an observer blinded to any preoperative and intraoperative Doppler-echocardiographic and surgical findings. After storage of 3D datasets, quantitative analysis was promptly performed. Three orthogonal mitral annulus images were displayed and subsequently modified to optimise visualisation of the entire annulus. Mitral annular measurements (annular area, circumference, intercommissural and anteroposterior diameter and height) were obtained in systole just before aortic valve closure. Annular height, maximal instantaneous vertical distance between highest (antero or posterior) and lowest (anterolateral or posteromedial) points, was used to compute the ratio of annular height to intercommissural diameter, a measure of annular saddle shape. After annular delineation, mitral leaflets were traced in multiple slices from medial to lateral commissure, and landmarks were placed at coaptation zone tips to define the entire surface of anterior and posterior leaflets. From valvular 3D data, mitral leaflet measurements (3D total valvular area, anterior leaflet area, posterior leaflet area) and quantification of leaflet excess movement in relation to annular plane (volume and height of prolapse) were also obtained at the same timing in systole (figure 3). Variability of measurements was determined by repeating measurements on stored 3D datasets at least 3 weeks after initial measurement by the same observer (intraobserver) and a different observer (interobserver) in 10 randomly selected cases.

**Clinical variables**

Baseline clinical data collected at preoperative echocardiogram and clinical visits included age, gender, weight, height, body mass index, body surface area, functional status and cardiac rhythm. Cardiovascular history and medications were determined by reviewing all medical records. Surgical variables were abstracted from surgical report dictated immediately after surgery completion, regarding CLI presence and extent (leaflets, segments) of mitral prolapse.

**Statistical analysis**

Data are shown as mean±SD for continuous variables and percentages for categorical variables. Patients were categorised according to 3DTOC presence/absence of posterior leaflet CLI and compared using t tests and χ² or Fisher exact tests as appropriate. For CLI diagnosis, the surgeon visual inspection (blinded to 3DTOC findings) was the diagnosis standard. For reproducibility, intraobserver and interobserver variability were calculated.

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**Figure 1** Example of a patient with myxomatous mitral valve disease (MMVD) and CLI with recording of 3D imaging and operative view imaging. Left image: 3D transoesophageal echocardiographic view of the mitral valve from the left atrial position. Right image: direct mitral valve view during surgical inspection. The red arrow indicates the CLI of the posterior mitral valve leaflet; the yellow arrow head indicates the prolapsing scallop (middle) of the posterior leaflet. Note, the visible deep indentation between P2 and P3 by both 3D imaging and direct viewing of the mitral valve. CLI, cleft-like indentation.
by dividing absolute differences between two measurements by their average value. All tests were two tailed; p<0.05 was considered significant. Analyses were performed using JMP V9.0.1 (SAS Institute, Cary, North Carolina, USA).

RESULTS
Clinical and surgical baseline characteristics are indicated in table 1. Overall, patients presented in the typical early sixth decade with male predominance. Risk factors were typical of age with generally normal renal function. For MR clinical manifestations, severe symptoms and atrial fibrillation were rare. Echocardiographic characteristics are presented in table 2, showing moderate LV dilatation consistent with MR associated with preserved LV function. Quantitative measurement of ERO, RVol and left atrial volume index clearly demonstrated that MR due to MMVD was severe. Bileaflet prolapse was noted in a minority of patients.

Among our 49 patients, none was diagnosed with CLI by 2D echo and 17 (35%) had CLI of posterior leaflet by 3DTOC. Of these, 15 were confirmed by the surgeon, and none was surgically diagnosed in patients without 3DTOC-determined CLI. Thus, for CLI diagnosis by 3DTOC, sensitivity was 100% and specificity 94%, positive predictive value 88% and negative predictive value 100%. 3DTOC location of CLI was unique in 15 patients (88%); in 8 (47%), CLI were located between P1–P2 scallops, and in 7 (41%) patients, CLI were located between P2–P3 scallops. Two patients had CLI of both interscallop indentations (between P1–P2 and P2–P3).

Stratified by 3DTOC-diagnosed CLI, there was no clinical difference between patients with and without CLI (table 1, right part), in particular with similar treatment in both groups proportionate to hypertension and symptoms. Among surgical variables, presence of flail segments was similar with or without CLI, but patients with CLI had more often single scallop posterior leaflet prolapse (82% vs 44%, p=0.007).

In regard to 2D Doppler echocardiography, comparison between those with and without CLI yielded limited differences (table 2, right part). Dimensions of LF, LA and severity of MR were similar with or without CLI. However, patients with CLI presented with slightly lower EF (60%±11% vs 65%±5%, p=0.03), although on average in the normal range. In all cases except one, mitral valve repair was successful (98%) and all patients with CLI diagnosed by the surgeon underwent closure of interscallop area during repair.

The 3DTOC quantified annular and valvular variables are presented in table 3. Overall, all dimensions of annulus in systole were very similar to previous studies of MMVD undergoing surgery for MR, which had demonstrated enlargement of all dimensions in comparison to normal subjects, with mild annular
flattening in systole. Despite this standard MMVD presentation and similar MR severity in both groups with similar LV and LA remodelling, there were multiple differences between patients with and without CLI by 3DTOC (table 3, right part). With CLI, the annulus was smaller for all systolic dimensions, anteroposterior (42.2±7.1 vs 47.0±7.5 mm, p=0.04), intercommissural (36.5±4.5 vs 41.4±5.9 mm, p=0.005), circumference (133±16 vs 148±19 mm, p=0.009) and area (1289±326 vs 1619±427 mm², p=0.008). There was only a trend for smaller annular height with CLI so that the ratio of annular height to intercommissural diameter, a measure of annular saddle shape, was not different with and without CLI. For valvular measurements, there were also marked 3DTOC differences between patients with and without CLI. There was a trend towards smaller volume of prolapse with CLI versus without CLI (1.9 ±1.2 vs 4.0±4.3 mL, p=0.06), which in view of similar maximum height of systolic prolapse suggests smaller prolapse base with CLI. Importantly, valve tissue was markedly smaller with versus without CLI overall (leaflets’ area 1574±409 vs 2019±652 mm², p=0.01), affecting both anterior and posterior leaflet (table 3). These differences are not linked to MR severity as 3D annular and valvular differences between patients with and without CLI were unchanged stratified by RVol (figure 4). Thus, CLI, as part of the phenotypical spectrum of MMVD, affect patients with single scallop prolapse, limited annular enlargement and relative tissue paucity, irrespective of MR severity.

Table 1  Clinical baseline characteristics and surgical data

<table>
<thead>
<tr>
<th></th>
<th>All patients, n=49</th>
<th>CLI absent, n=32</th>
<th>CLI present, n=17</th>
<th>p Value</th>
</tr>
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<tr>
<td>Age, years</td>
<td>63±11</td>
<td>64±10</td>
<td>60±14</td>
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<td>Male gender, %</td>
<td>78</td>
<td>78</td>
<td>76</td>
<td>0.89</td>
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<td>Body surface area, m²</td>
<td>1.97±0.23</td>
<td>1.98±0.22</td>
<td>1.95±0.26</td>
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<td>Body mass index, kg/m²</td>
<td>26±4</td>
<td>27±4</td>
<td>26±3</td>
<td>0.38</td>
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<tr>
<td>Dyslipidaemia, %</td>
<td>41</td>
<td>38</td>
<td>47</td>
<td>0.52</td>
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<tr>
<td>Hypertension, %</td>
<td>43</td>
<td>44</td>
<td>41</td>
<td>0.86</td>
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<tr>
<td>Diabetes, %</td>
<td>8</td>
<td>9</td>
<td>6</td>
<td>0.66</td>
</tr>
<tr>
<td>History of CAD, %</td>
<td>8</td>
<td>6</td>
<td>12</td>
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<tr>
<td>Atrial fibrillation, %</td>
<td>14</td>
<td>13</td>
<td>18</td>
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<tr>
<td>NYHA class, %</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>I</td>
<td>31</td>
<td>34</td>
<td>24</td>
<td>0.86</td>
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<tr>
<td>II</td>
<td>43</td>
<td>41</td>
<td>47</td>
<td>0.86</td>
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<tr>
<td>III</td>
<td>22</td>
<td>22</td>
<td>23</td>
<td>0.73</td>
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<td>IV</td>
<td>4</td>
<td>3</td>
<td>6</td>
<td>0.90</td>
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<td>Creatinine, mg/dL</td>
<td>1.0±0.2</td>
<td>1.0±0.2</td>
<td>1.0±0.2</td>
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<td>Medical therapy, %</td>
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<td></td>
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<tr>
<td>ACE inhibitors/ARBs</td>
<td>43</td>
<td>44</td>
<td>41</td>
<td>0.03</td>
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<tr>
<td>β-blockers</td>
<td>33</td>
<td>34</td>
<td>29</td>
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<td>Diuretics</td>
<td>24</td>
<td>25</td>
<td>24</td>
<td>0.73</td>
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<td>Surgical variables</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td>Bypass time, min</td>
<td>72±29</td>
<td>71±30</td>
<td>74±28</td>
<td>0.72</td>
</tr>
<tr>
<td>Flail leaflet, %</td>
<td>61</td>
<td>59</td>
<td>65</td>
<td>0.71</td>
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<tr>
<td>Posterior leaflet prolapsing segments</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Single scallop, %</td>
<td>57</td>
<td>44</td>
<td>82</td>
<td>0.007</td>
</tr>
<tr>
<td>Multiple scallops, %</td>
<td>43</td>
<td>56</td>
<td>18</td>
<td>0.04</td>
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<tr>
<td>Valve repair, %</td>
<td>98</td>
<td>97</td>
<td>100</td>
<td>0.35</td>
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</tbody>
</table>

ARF, angiotensin receptor blocker; CAD, coronary artery disease; CLI, cleft-like indentation; NYHA class, New York Heart Association functional class.

Table 2  2D echocardiographic baseline characteristics

<table>
<thead>
<tr>
<th></th>
<th>All patients, n=49</th>
<th>CLI absent, n=32</th>
<th>CLI present, n=17</th>
<th>p Value</th>
</tr>
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<tr>
<td>LVEF, %</td>
<td>63±7</td>
<td>65±5</td>
<td>60±11</td>
<td>0.03</td>
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<td>LVEDD, mm</td>
<td>57±7</td>
<td>57±6</td>
<td>57±6</td>
<td>0.91</td>
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<td>LVESD, mm</td>
<td>36±5</td>
<td>36±4</td>
<td>37±7</td>
<td>0.34</td>
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<tr>
<td>LAVI, mL/m²</td>
<td>60±17</td>
<td>62±16</td>
<td>59±20</td>
<td>0.55</td>
</tr>
<tr>
<td>ERO, cm²</td>
<td>0.50±0.23</td>
<td>0.49±0.21</td>
<td>0.54±0.24</td>
<td>0.54</td>
</tr>
<tr>
<td>RVol, mL/beat</td>
<td>85±48</td>
<td>89±56</td>
<td>79±29</td>
<td>0.49</td>
</tr>
<tr>
<td>SPAP, mm Hg</td>
<td>35±12</td>
<td>35±9</td>
<td>35±15</td>
<td>0.98</td>
</tr>
<tr>
<td>Single-leaflet MMVD</td>
<td>63% (31)</td>
<td>59% (19)</td>
<td>71% (12)</td>
<td>0.43</td>
</tr>
<tr>
<td>Bileaflet MMVD</td>
<td>37% (18)</td>
<td>41% (13)</td>
<td>29% (5)</td>
<td>0.43</td>
</tr>
</tbody>
</table>

CLI, cleft-like indentation; EDD, end-diastolic diameter; ERO, effective regurgitant orifice area; ESD, end-systolic diameter; LAVI, left atrial volume index; MMVD, myxomatous mitral valve disease; RVol, regurgitant volume; SPAP, systolic pulmonary artery pressure.
Intraobserver and interobserver variability for 3DTOC measurements were small (AP diameter 3.3%±1.1% and 5.1% ±3.7%, annular height 4.6%±0.3% and 1.4%±0.6%, annular circumference 2.2%±2.8% and 2.3%±6.4%, valvular area 3.3%±0.3% and 7.4%±0.8%) and were similar to those previously reported.17

**DISCUSSION**

The present study reports on diagnosis and morphological associations of CLI of mitral valve in patients with MMVD undergoing mitral surgery for severe MR. Our study shows that (1) diagnosis of posterior leaflet CLI, which was not feasible by standard 2D echocardiography, was reliable by 3DTOC
with high sensitivity and specificity; (2) CLI are not an uncommon finding (one-third of patients) in MMVD and requires specific treatment during valve repair and (3) mechanistically, CLI are not explained by excessive annular enlargement that would cause mitral valve stretching or by excessive leaflet prolapse that would cause excess scallop separation. Hence, CLI are part of a global morphological spectrum within MMVD, occurring in the context of single scallop prolapse with insufficient leaflet tissue to cover the marked indentation separating prolapsing and non-prolapsing segments. This wide separation may be accentuated by leaflet resorption during valve repair, which underscores the importance of diagnosing CLI by 3DTOC prebypass to conduct proper heart-team communication and valve repair.

Importance of CLI in MMVD

In the general adult population, CLI of posterior mitral leaflet are rare, with very low prevalence (~0.1%) in one echocardiographic study, but may be slightly higher in normal controls.12 CLI have been mentioned in Marfan syndrome19 with associated anomalies of the mitral apparatus, such as malrotation of papillary muscles.20 In patients with MR, CLI were noted to occur more frequently in MMVD than in patients with other causes of MR.12 The quantitative criteria for defining CLI versus normal indentations between posterior mitral scallops remain tentative, but we are confident that using the previously suggested 50% minimum of adjacent scallops’ depth as diagnostic criterion for CLI12 16 is concordant with surgical definition of CLI. However, because CLI are part of MMVD morphological spectrum, as shown by our study, CLI prevalence among MMVD populations may depend on the subset enrolled (single vs multi-scallop MMVD in particular) possibly explaining prevalence variability. Careful detailed examination of the entire posterior leaflet by 3DTOC is the key to diagnosis, but future collaborative studies are warranted to develop quantitative 3D criteria for CLI diagnosis now that appropriate imaging and software allow in-depth analysis of mitral complex.17 21

Study of valve characteristics such as CLI has been affected by confusion attached to the label ‘cleft’. The typical cleft is a gap at the middle of anterior leaflet, but it has been long known that rarely posterior leaflet may be affected by such true cleft involving central tissue gap.9 Diagnosis of this typical pattern of congenital cleft mitral valve was recently enhanced by 3DTOC.10 11 22 However, use of the term ‘cleft’ in describing indentations between posterior leaflet scallops, normal or exaggerated,14 has been confusing. Hence, we believe, it is essential to separate nosologically CLI from true clefts affecting the posterior leaflet.9 11 12 In terms of diagnosis, 2DTEE may have difficulty ascertaining typical cleft despite multiplane imaging23 24 with even more difficulty with CLI, so that CLI identification is essential in demonstrating 3DTOC usefulness.5 Multiple jets and gaping separations between scallops may be clues to CLI, but definite diagnosis requires 3DTOC. Hence, preoperative identification of CLI may be the first strong indication that 3DTOC is important and irreplaceable by 2DTEE. CLI identification takes all its importance in view of recent guidelines that place high-quality repair as key to the surgical indication in degenerative MR.3 3DTOC, by identifying in systole CLI of posterior leaflet while the heart is loaded, has definitive advantages over flaccid heart examination and allows quantitative assessment of valvular tissue morphology to define its mechanism. However, 3D pitfalls in interpreting leaflet gaps, such as echo dropout (annular calcification, improper gain) or oblique imaging planes creating the impression of a fissure not actually present, emphasise proper en-face image acquisition and interpretation for 3DTOC.

Mechanisms of CLI by 3DTOC

Our working hypothesis that marked annular enlargement may lead to excessive separation of scallops was not verified. The rationale for this hypothesis was previous identification of asymmetric mitral annular enlargement in MMVD, sparing the inter-trigonal component (anterior) and dilating the remaining annulus essentially posterior.23 However, no measure of annular dimension was larger in patients with CLI, definitely ruling out annular enlargement as potential mechanism. A secondary hypothesis of excess prolapse causing CLI is also not verified as depth and volume of prolapse were not higher in patients with CLI.12 Conversely, our study suggests that CLI occur in a specific morphological context, involving single scallop prolapse of posterior leaflet and involving tissue paucity in contrast to patients without CLI. The tissue amount difference between CLI and non-CLI was not limited to posterior leaflet so that overall CLI were not detectable with marked excess global tissue. The prominent indentation may be conformational with native valve tissue paucity or acquired in patients whose response does not involve marked tissue redundancy despite MMVD and severe MR. Indeed, some valve segments may appear normal with isolated severe prolapse or flail segment, usually involving the middle scallop of posterior leaflet.26 Acquired accentuation of indentations is also possible, whereby normal positioning of two of the three scallops of posterior leaflet may strain their confluence with the single prolapsing scallop. Normal indentations act much the way pleats do on clothing, allowing leaflets to accommodate the curved shape of valve closure line.27 In contrast to patients with CLI, those with multiple prolapsing scallops incur less stretching of normal indentations. Hence, 3DTOC provides mechanistic insights that CLI are associated mostly with single scallop prolapse and insufficient tissue to cover the scallop separation, probably combining conformational and acquired valvular patterns. This data further emphasises the complex combination of various morphological elements in MMVD21 and warrants detailed physiological future studies of annular and valvular dynamics with quantitative MR dynamic assessment to characterise the complete phenotypical manifestations of MMVD.

Practical implication

Accurate characterisation of MMVD lesion’s complexity is critically important to predict feasibility of mitral valve repair and prognosis.28 This is increasingly relevant given the growing referral of asymptomatic patients for mitral valve surgery in whom predictable repair avoiding recurrent MR is crucial.3 MR of MMVD may be caused by lesions’ combination, to which CLI may contribute through extension of the valvar gap, and its satisfactory correction requires addressing all lesions at surgery.29 This is particularly important because valvular resection of posterior leaflet middle scallop may accentuate indentation width and cause residual MR and implies careful closure by the surgeon.30 Therefore, identification of CLI should be diligent8 and may also be important with the development of catheter-based MR-correction techniques.31 In that regard, future resolution improvements of colour-flow imaging may contribute to CLI diagnosis and to the interpretation of its role as a cause of MR.

Limitations of the study

Population size may have limited our ability to detect borderline differences such as prolapse volume. However, major differences between patients with and without CLI are clearly established...
by our mechanistic 3DTOC study. Larger samples will be necessary in verifying CLI diagnostic accuracy of 3DTOC in clinical practice and the implications of CLI suturing in preventing recurrent MR postrepair. In this study, we could not determine with certainty whether MR occurred through the CLI. With the continued progress of colour-flow imaging in 3DTOC, future studies should focus on defining CLI contribution to lateral or multiple MR jets.

CONCLUSION

CLI of posterior leaflet are frequent in MMVD, require closure during mitral repair and are identified by 3DTOC with high accuracy. CLI are mechanistically unrelated to excess annular enlargement or excess prolapse, but conversely occur when single scallop prolapse/flail is associated with relative tissue paucity. These 3DTOC data enhance diagnostic and mechanistic comprehension of the diversity of MMVD phenotypical presentation.

Key messages

What is already known on this subject?

Recently described clef-like indentations (CLI) of posterior mitral leaflet in myxomatous mitral valve disease (MMVD) can be associated with mitral regurgitation (MR) and failure of mitral repair.

What might this study add?

CLI diagnosis is not feasible by standard 2D echocardiography, but is reliable by 3D transesophageal echocardiography (3DTOC) and requires specific treatment during valve repair. Mechanistically, CLI are not explained by excessive annular enlargement or excessive leaflet prolapse, but rather by single scallop prolapse and tissue paucity causing excess posterior scallops separation.

How might this impact on clinical practice?

Accurate characterisation of MMVD lesion’s complexity—essential to define mitral repair feasibility and success—underscores the importance of 3DTOC and potentially its timing. 3DTOC diagnosis of CLI warrants consideration of CLI closure. The finding that tissue paucity is linked to CLI may influence the choice and magnitude of tissue ressection during mitral valve repair.

Contributors All authors have contributed to the concept of the study, data collection, data analysis and interpretation, drafting or revision of the manuscript.

Competing interests None declared.

Ethics approval Mayo Clinic Institutional Review Board.

Provenance and peer review Not commissioned; externally peer reviewed.

REFERENCES


