Importance of Mitral Subvalvular Apparatus in Terms of Cardiac Energetics and Systolic Mechanics in the Ejecting Canine Heart

Kwok L. Yun, Marek A. Niczyporuk, George E. Sarris, James F. Fann, and D. Craig Miller
With the technical assistance of Geraldine C. Derby and Cynthia E. Handen
Department of Cardiovascular Surgery, Stanford University School of Medicine, Stanford, California 94305

Abstract
To assess the importance of the intact mitral subvalvular apparatus for left ventricular (LV) energetics, data from nine open-chest ejecting canine hearts were analyzed using piezoelectric crystals to measure LV volume. After mitral valve replacement with preservation of all chordae tendineae, baseline LV function was assessed during transient caval occlusion: A quadratic fit of the LV end-systolic pressure–volume data was used to determine the curvilinear end-systolic pressure–volume relationship (ESPVR). All chordae were then divided with exteriorized snares. Reassessment revealed deterioration of global LV pump function: (a) the coefficient of nonlinearity, decreased (less negative) by 90% (P = 0.06); (b) slope of the curvilinear ESPVR at the volume axis intercept, decreased by 75% (P = 0.01); and V_{100}, end-systolic volume at 100 mmHg end-systolic pressure, increased by 42% (P < 0.02). Similarly, preload recruitable stroke work fell significantly (−14%) and V_{1,000} (end-diastolic volume [EDV] at stroke work [SW] of 1,000 mmHg · ml) rose by 17% (P < 0.04). With respect to LV energetics, the total mechanical energy generated by the ventricle decreased, as indicated by a decline in the slope of the pressure volume area (PVA)-EDV relationship (120±13 [mean±SD] vs. 105±13 mmHg, P < 0.001). Additionally, comparison of LV SW and PVA from single beats with matched EDV showed that the efficiency of converting mechanical energy to external work (SW/PVA) declined by 14% (0.65±0.13 vs. 0.56±0.08, P < 0.03) after chordal division. While effective systemic arterial elastance, E_{as}, also fell significantly (P = 0.03) after the chordae were severed, the E_{as}/E_{es} ratio (E_{es} = slope of the linear ESPVR) increased by 124% (0.91±0.53 vs. 2.04±0.87, P = 0.001) due to a proportionally greater decline in E_{es}. This indicates a mismatch in ventricle–arterial interaction, deviating from that required for maximal external output (viz., E_{as}/E_{es} = 1). These adverse effects of chordal division may be related to the observed changes in LV geometry (i.e., eccentricity). We conclude that the intact mitral subvalvular apparatus is important in optimizing LV energetics and ventriculo–vascular coupling in addition to the enhancement of LV systolic performance. (J. Clin. Invest. 1991. 87:247–254.) Key words: chordae tendineae • LV function • ventriculo–vascular interaction

Introduction
Mitral papillary-annular continuity has been hypothesized to be important for optimal global left ventricular (LV) systolic performance, a concept we have previously termed “valvular–ventricular interaction” (1). In isovolumic experimental preparations, it has been shown that chordal division results in deterioration of LV systolic mechanics (1–6), which can be restored by reattachment of the mitral chordae (5). The anterior and posterior leaflet chordae have similar, but additive, contributions to LV systolic elastance (4). More recently, the importance of the integrity of the mitral subvalvular apparatus for LV pump function has been confirmed in in situ ejecting canine hearts using relatively load-insensitive indexes of LV systolic function (7); chordal severing resulted in a 72% decline in E_{as}, slope of the linear end-systolic pressure–volume relationship (ESPVR). The extrapolated volume intercept (V_{0}) also fell significantly, but was in the (nonphysiologic) negative range (−16.6±15.3 ml) after chordal disruption. The possibility that ESPVR was nonlinear outside the physiologic pressure range examined or whether there was an actual change in the curvilinearity of the ESPVR is unknown. Moreover, the implications of this deterioration in systolic mechanics in terms of LV energetics and efficiency were not examined in the setting of altered LV geometry (vs. change in contractile state). We therefore assessed the effects of chordal division on LV myocardial energetics, mechanical energy (pressure volume area) to external pressure–volume work efficiency, and systolic pump function using curvilinear ESPVR analysis. It is important that ventriculo–arterial coupling, which ultimately determines the efficiency of energy transfer to the systemic circulation, was examined before and after chordal severing, since LV systolic function may not necessarily be related to ventriculo–vascular interaction (8).

Methods
Surgical preparation. This surgical preparation has been described in detail previously (7). Briefly, nine adult, mongrel dogs (25–30 kg) were anesthetized with thiopental (25 mg/kg) and maintained with 0.5–1%

1. Abbreviations used in this paper: E_{as}, effective systemic arterial elastance; E_{es}, slope of the linear ESPVR; EDD, end-diastolic volume; ESPVR, end-diastolic pressure; EDV, end-diastolic volume; EED, end-ejection diameter; EM, electromagnetic; EF_PVR, end-systolic pressure–volume relationship; LV, left ventricular; MEP, mean ejection pressure; %ID, fractional diameter shortening; P_{as}, end-systolic pressure; PRSW, preload recruitable stroke work; PVA, pressure–volume area; SW; stroke volume; SV_{es}, SV as determined by sonomicrometry; SW; stroke work; V_{as}, end-systolic volume; V_{es}, volume axis intercept; V_{pva}, preload recruitable PVA volume intercept.
halothen. The heart was exposed via a left lateral thoracotomy, and sets of piezoelectric crystals were implanted to allow continuous measurement of three orthogonal LV dimensions (base-apex [Dba], anterior-posterior [Dap], and septal-lateral [Dsl]). A siltastic occluder was placed around the inferior vena cava to alter preload abruptly. On cardiopulmonary bypass the mitral valve was exposed through a left atriotomy, and all chordae were snared with 0-Flaxen wire sutures and exteriorized through plastic tubing that was secured to the epicardium. A 25-mm porcine bioprosthetic valve was then sutured to the mitral annulus with complete preservation of the subvalvular apparatus. The atriotomy was closed, and the animal heaned from cardiopulmonary bypass. A micrometer-tipped catheter was inserted into the LV cavity via the apex, and a fluid-filled catheter was inserted into the left atrium to guide volume replacement. An electromagnetic (EM) flow probe was placed around the ascending aorta to measure instantaneous flow. This report is based on further analysis of data derived from the same animals reported earlier by Sarris et al. (7).

**Experimental protocol.** After 1 h of reperfusion and stabilization of arterial pressure and cardiac output, baseline hemodynamics and dimensional data were obtained over a physiologic range of peak systolic LV pressures by transient inferior vena cava occlusion (during which the ventilator was temporarily disconnected to minimize respiratory variation). Three baseline measurements were obtained, allowing 3–5 min between data acquisition runs for steady-state equilibration. After baseline data were collected, all chordae tendineae were divided by pulling the wire suture snare, and data again collected in triplicate. Warm, fresh, homologous blood was infused to maintain baseline levels of mean arterial pressure.

Analogue data were recorded on a multichannel recorder (model 7758B; Hewlett-Packard Co., Andover, MA) at a paper speed of 25 mm/s. The following parameters were measured: LV pressure (P), electronically differentiated LV dp/dt, ECG, aortic EM flow, and the three dimensions Dba, Dap, and Dsl. These data were subsequently digitized at 250 Hz by a 6942A multiprogrammer (Hewlett-Packard), controlled by a computer workstation (series 9000, model 216; Hewlett-Packard), and stored on digital disk drives.

**Data analysis.** The data were analyzed on a series 9000, models 226 and 310 computer workstations using software developed in our laboratory. Premature ventricular beats and the subsequent two to four extra-systolic beats were excluded from analysis by an R-R interval sampling algorithm. Instantaneous LV volume V(t) was calculated by assuming the ventricle to be an ellipsoid and subtracting the directly measured (postmortem) ventricular muscle volume (Vmv(t)) (7): V(t) = (π/6) (Dba x Dap x Dsl) - Vmv(t). The following variables were calculated for each cardiac cycle: End-diastolic volume (EDV) and end-ejection volume (EEV) (defined as LV volume at peak +dp/dt and peak -dp/dt, respectively), stroke volume (SV) (computed as EDV - EEV), cardiac output (CO) (calculated as SV x heart rate), mean ejection pressure (MEP) (determined as the time integral of LV pressure during ejection defined by the aortic flow signal), and total peripheral resistance (estimated as MEP/CO). Stroke work (SW) was computed as the time integral of LV pressure and volume for each cardiac cycle: SW = \int P x dV. Since the estimation of LV volume is dependent on chamber geometry, SV derived from sonomicrometric measured LV dimensions for each beat during caval occlusion was compared with that obtained from the time integral of the aortic EM flow signal during ejection, both before and after chordal division, to assure that chordal severance did not alter the relation between true LV volume and that estimated using the ellipsoid model.

End-systolic pressure (Pse) and volume (Vse) were determined using an iterative technique (9, 10). To avoid problems with linear extrapolation of a curvilinear relation beyond the data range (11, 12), LV volume at an end-systolic pressure of 100 mmHg, Vse, was calculated. Alternatively, end-systolic data were also fitted to the following parabolic curvilinear model (12): Pse = a (Vse - V0)^2 + b (Vse - V0) + c. In this expression, a is the coefficient of curvilinarity, V0 is the volume axis intercept, and b is the slope of the ESPVR at V0 using a best least-squares algorithm.

Another load-insensitive measure of contractile state, preload recruitable SW (PRSW), was determined during each caval occlusion by least-squares linear regression relating SW to EDV (13): SW = Msw (EDV - Vse), where Msw is the slope and Vse is the volume intercept. Again, to avoid extrapolation beyond the range of the observed data, EDV at an SW of 1,000 mmHg x mL, Vse, was calculated in addition to Vse.

The pressure volume area (PVA) (represented as the total area under the nonlinear ESPVR as well as the systolic segment of the pressure–volume trajectory and above the end-diastolic pressure–volume relation curve) was calculated for each beat during caval occlusion by the method of Goto et al. (14). Preload recruitable PVA (PRPVA) was defined as the slope of the least-squares linear regression between PVA and EDV: PVA = Mpv (EDV - Vpv), where Mpv and Vpv are the slope and volume intercept, respectively. The efficiency of energy transfer from PVA to external mechanical pressure–volume work (SW/PVA) (15) was calculated during each data acquisition run for beats with matched LV EDV. Effective arterial elastance (Ea) was defined as Ppv / (EDV - Vpv) by Sunagawa (16), was calculated from the first beat of each caval occlusion during steady-state conditions. The Ea/Epv ratio was then computed, where Epa represents the slope of the linear ESPVR.

In the study of effect of chordal division on LV chamber geometry, the following parameters were calculated for each of the three LV axes studied at a matched LV end-diastolic pressure (EDP): End-diastolic diameter (EDD) and end-ejection diameter (EED) were defined at the time of EDV and EEE, respectively. Percent systolic fractional shortening (%ES) was calculated as [(EED - EEDV)/EED] x 100. Eccentricity (e) was computed as (17): e = \sqrt{Dba^2 - Dap^2/Dba^2}.

**Statistical analysis.** All results are expressed as mean±1 SD unless specified otherwise. Mean values for all variables, calculated as the average of all three measurements at baseline and after chordal division, were compared with a two-tailed, paired t test. Comparisons of correlation coefficients were made after Fisher’s z-transformation. The level of significance was taken as P ≤ 0.05. A P value > 0.05 but ≤ 0.10 was considered to be possibly significant and is stated as such.

**Results**

**Hemodynamics.** Table I summarizes the effects of chordal division on LV hemodynamic parameters. As previously reported (7), dividing the chordae resulted in a small but significant decline in heart rate (from 136±30 to 130±27 beats/min, P = 0.04). Despite an increase in EDV (from 34±14 to 40±15 ml, P = 0.05), SV and CO did not change. Compared with baseline values, MEP fell significantly from 107±11 to 83±19 mmHg (P = 0.02) after chordal severing, resulting in a possibly significant decline (P = 0.09) in calculated total peripheral resistance. Moreover, the relation between SV as determined by sonomicrometry (SVsono) and aortic EM flow (SV preferring) did not change after severing the chordae tendineae, as illustrated in Fig. 1. At baseline, SVsono was linearly related to SV preferring by the equation: SVsono = (0.65±0.09) x (SV preferring) + (10.17±2.70), with r ranging from 0.956 to 0.998. After chordal division, this relation remained linear (r = 0.967-0.994) without any significant alteration in either the slope (0.67±0.06) or the x-axis intercept (0.34±5.02 ml).

**Global left ventricular systolic performance.** Fig. 2 graphically illustrates the effects of chordal division on the nonlinear ESPVR; data for all animals are summarized in Fig. 3. The nonlinear model fit the observed ESPVR data closely throughout the LV pressure range, with nonlinear correlation coefficients ranging from 0.962 to 0.998 and 0.943 to 0.998 before and after chordal severing, respectively (P = NS). Although the second order coefficient a varied markedly, the overall value of

K. L. Yun, M. A. Niczyproruk, G. E. Sarris, J. I. Fann, and D. C. Miller
Table I. Hemodynamic Parameters before and after Chordal Division

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<tr>
<td>Heart rate (min⁻¹)</td>
<td>136±30</td>
<td>130±27*</td>
</tr>
<tr>
<td>LV end-diastolic volume (ml)</td>
<td>34±14</td>
<td>40±15*</td>
</tr>
<tr>
<td>LV stroke volume (ml)</td>
<td>23±9</td>
<td>25±9</td>
</tr>
<tr>
<td>Cardiac output (litter/min)</td>
<td>3.2±1.4</td>
<td>3.2±1.4</td>
</tr>
<tr>
<td>Total peripheral resistance (mmHg·s/ml)</td>
<td>2.52±1.50</td>
<td>2.10±1.38*</td>
</tr>
<tr>
<td>Mean LV ejection pressure (mmHg)</td>
<td>107±11</td>
<td>83±19*</td>
</tr>
</tbody>
</table>

Results are expressed as mean±1 SD.
Vs. chordae intact: * P < 0.05; † P = 0.09; ‡ P = 0.02

a for all runs was −3.149±5.755 mmHg/ml² (95% confidence limits: −6.216 and −0.082 mmHg/ml²); therefore, the ESPVR was significantly nonlinear, being concave towards the abcissa (LV volume axis). After division of the subvalvular apparatus, the ESPVR became less concave, but this difference was only possibly significant (a = −5.7±17.4 vs. −0.6±0.8 mmHg/ml²; P = 0.064). V₀ fell (8.8±5.7 vs. 3.4±7.0 ml; P = 0.009), but remained within the physiologic range. Systolic function, however, as estimated by the slope of the nonlinear ESPVR at V₀, b, declined markedly by 75% after chordal division (46±34 vs. 12±8 mmHg/ml; P = 0.01). V₁₀₀ increased significantly by 42% from 17±10 to 25±15 ml (P = 0.014) after the chordae were severed.

As a corollary, changes in PRSW (another load-insensitive measure of LV systolic performance which incorporates both systolic and diastolic functions [12]), were also analyzed (Fig. 3). The SW–EDV relationship was linear both before (r = 0.959–0.997) and after chordal severing (r = 0.976–0.998).

After the chordae were divided, Mₑ declined by 14% (81±13 vs. 69±16 mmHg; P = 0.02) without consistent changes in the volume intercept, Vₑ (13.3±7.9 vs. 16.2±7.2 ml; P = 0.14). As was seen for V₁₀₀, Vₑ1,0₀ increased by 17% from 27±8 to 32±9 ml (P = 0.037).

**Left ventricular energetics.** Fig. 4 illustrates the effect of chordal severing on PRPVA as defined above. In this representative animal, the slope, Mₑ, fell from 105 to 84 mmHg with minimal reduction in the volume intercept, Vₑ. In all dogs, the PVA–EDV relationship was highly linear at baseline (r ranged from 0.990 to 0.999) and was not affected by chordal severing (r = 0.995–1.00; P = NS). Comparison of measurements after chordal division with baseline values for all animals revealed a highly significant 13% decline in Mₑ (120±13 vs. 105±13 mmHg; P < 0.001) and a 9% decrease in Vₑ (15.0±6.6 vs. 13.7±6.7 ml; P = 0.033). This implied a reduction in total mechanical work able to be generated at any given LV EDV. Furthermore, the SW/PVA ratio at matched EDV, reflecting the efficiency of energy transfer from PVA to external mechanical pressure–volume work, also decreased significantly by 14% (0.65±0.13 vs. 0.56±0.08; P = 0.023) after chordal disruption (Fig. 5).

When compared with baseline values, steady-state effective arterial elastance, Eₑa, fell significantly by 43% from 8.6±5.1 to 4.9±1.5 mmHg/ml (P = 0.031) with chordal severing. In terms of ventriculo–arterial coupling, however, the Eₑa/Eₑa ratio (Eₑa obtained using linear regression within only a limited LV pressure range) actually increased by over twofold (0.91±0.53 vs. 2.04±0.87; P = 0.001), because of a proportionally greater (−72%) decline in Eₑa (Fig. 6). With the chordae intact, the 95% confidence intervals for Eₑa/Eₑa ranged from 0.47 to 1.26, which were not significantly different from unity.

The effects of chordal severing on LV chamber geometry at matched LV EDV are summarized in Table II. After chordal division, the increase in EDV was due to some lengthening of EDD in all three LV axes, but primarily along the major axis. In contrast, %D rose in the anterior-posterior (P < 0.05) and possibly the septal-lateral (P = 0.054) dimensions. Fractional diameter change during ejection in the base-apex direction was unaltered due to a parallel increase in EED (P < 0.001). As a result, the LV was significantly more eccentric at end-diastole; this was possibly also true at end-ejection (P = 0.067).

**Discussion**

The concept valvular-ventricular interaction was recently demonstrated in in situ ejecting canine hearts by Sarris et al. (7).
The integrity of the mitral subvalvular apparatus significantly influenced global LV systolic performance as reflected by a 72% decline in $E_{es}$, the slope of the linear ESPVR, after chordal severing. However, the extrapolated volume axis intercept decreased significantly into the (nonphysiologic) negative range. Reduced $V_0$, in theory, would suggest an increase in systolic function as a totally unloaded ventricle can shorten more when pump performance is enhanced; moreover, the linearly extrapolated $V_0$ tends to change in the same direction as $E_{es}$ in response to inotropic interventions (18, 19). The data, therefore, can be interpreted as either nonlinearity of the ESPVR at high and low contractile states (12) or a change in the curvilinearity of the ESPVR, viz., a function of inotropic state (11, 12). In this study, a quadratic equation provided a significantly better fit of the end-systolic pressure volume points than did linear regression since the coefficient $a$ overall was significantly less than zero. After chordal division, $a$ was smaller (less negative) in all animals suggesting reduced concavity of the ESPVR with respect to the volume axis. This change attained only possible statistical significance, however, probably due to the high variability between individual dogs. $V_0$ was significantly lower, however, again reflecting the decrease in concavity of the curvilinear ESPVR with respect to the volume axis. The range of $V_0$, however, remained well within the physiologic range (vs. the $V_0$ values derived by linear regression [7]), supporting the appro-

**Figure 3.** Summary of the effects of chordal division on parameters characterizing nonlinear ESPVR and PRSW. Error bars represent±1 standard deviation. (a) Coefficient of curvilinearity; (b) slope of ESPVR at $V_0$; $V_0$, volume axis intercept of the nonlinear ESPVR; $V_{100}$, end-systolic volume at end-systolic pressure of 100 mmHg. $M_w$, slope of the stroke work–end-diastolic pressure relationship; $V_{es}$, PRSW end-diastolic volume axis intercept; $V_{1000}$, PRSW end-diastolic volume at stroke work of 1,000 mmHg·ml.

**Figure 4.** Representative pressure volume area–end-diastolic volume (PVA–EDV) relations obtained during caval occlusion before (closed circles) and after (open circles) chordal division, which were linear within the physiologic pressure range. In each animal, there was a decline in the slope of the PVA–EDV relationship, indicating reduced mechanical energy generated at an equivalent preload.
The method avoids extrapolation beyond the observed data range. Moreover, a similar value is calculated independent of the degree of polynomial fit (linear vs. quadratic). In this study, chordal division was associated with a 42% increase \( (P = 0.014) \) in \( V_{100} \), indicating a larger \( V_m \) at similar \( P_a \): this shift of the linear ESPVR to the right suggests a reduction in LV systolic mechanics.

Because the coefficients of the quadratic fit are not entirely orthogonal (some degree of interdependence is present), PRSW was also examined for corroborative evidence. The SW-EDV

![Graph of Figure 6](image)

**Figure 6.** The effects of chordal division on \( E_a \) and the \( E_a/E_m \) ratio. The increase in the \( E_a/E_m \) ratio indicates a mismatch between the left ventricle and the systemic arterial load. Error bars represent ±1 SEM.

<table>
<thead>
<tr>
<th>Chordal status</th>
<th>Intact</th>
<th>Divided</th>
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<tbody>
<tr>
<td>Base-apex axis ( (D_{BA}) )</td>
<td>81.6±5.4</td>
<td>84.8±5.4*</td>
</tr>
<tr>
<td>EDD (mm)</td>
<td>81.6±5.4</td>
<td>84.8±5.4*</td>
</tr>
<tr>
<td>EED (mm)</td>
<td>81.6±5.4</td>
<td>84.8±5.4*</td>
</tr>
<tr>
<td>dD (%)</td>
<td>-0.1±2.9</td>
<td>-0.1±3.4</td>
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<tr>
<td>Anterior-posterior axis ( (D_{AP}) )</td>
<td>59.0±5.2</td>
<td>60.1±4.6*</td>
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<tr>
<td>EDD (mm)</td>
<td>59.0±5.2</td>
<td>60.1±4.6*</td>
</tr>
<tr>
<td>EED (mm)</td>
<td>59.0±5.2</td>
<td>60.1±4.6*</td>
</tr>
<tr>
<td>dD (%)</td>
<td>3.2±3.3</td>
<td>4.7±2.7*</td>
</tr>
<tr>
<td>Septal-lateral axis ( (D_{SL}) )</td>
<td>45.7±3.8</td>
<td>47.4±3.8*</td>
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<tr>
<td>EDD (mm)</td>
<td>45.7±3.8</td>
<td>47.4±3.8*</td>
</tr>
<tr>
<td>EED (mm)</td>
<td>45.7±3.8</td>
<td>47.4±3.8*</td>
</tr>
<tr>
<td>dD (%)</td>
<td>10.9±5.0</td>
<td>13.8±5.0*</td>
</tr>
<tr>
<td>End-diastolic eccentricity index</td>
<td>0.687±0.051</td>
<td>0.701±0.051*</td>
</tr>
<tr>
<td>End-ejection eccentricity index</td>
<td>0.718±0.017</td>
<td>0.733±0.049*</td>
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Vs. chordae intact: * \( P < 0.001; \) * \( P = 0.06; \) * \( P < 0.05; \) * \( P = 0.054; \) * \( P = 0.067.\)
relation was highly linear; after chordal severing, the slope ($M_a$) decreased without any significant change in the volume axis intercept, indicating reduced global pump function. It should be noted, however, that $M_a$ fell by only 14% compared with 39% reported earlier (7). This discordance may be explained by the different method of SW calculation: SW had been calculated from aortic flow measurements in our previous study by the equation: $SW = SV \cdot (MEP – EDP)$. Such external work based on aortic flow or cardiac output, however, does not fully describe the mechanical work performed by the ventricle since diastolic work may be inaccurately determined, particularly at high filling pressures. Given the absence of associated alterations in the end-diastolic pressure–volume relation after chordal division (7), the increase in $EDV$ would imply that the ventricle was operating in a higher range of end-diastolic pressures, thereby resulting in greater underestimation of SW after chordal division. This may, in part, be responsible for the relatively greater reduction in PRSW when aortic flow was used to compute SW (7). Despite widespread variation between animals, $V_{a,1.00}$ increased significantly by 17% after chordal division, indicating a greater preload for equivalent external work; this rightward shift of the SW–EDV relation also implies deterioration in LV systolic pump function.

In the time-varying elastance model of the left ventricle, PVA represents the total mechanical energy generated by each contraction (24–26). It consists of external mechanical work performed during systole and elastic potential energy stored in the myocardium at end-systole. Like PRSW, the PVA–EDV relation was highly linear in this current experiment, and was unaffected by chordal severing. After disruption of the subvalvular apparatus, the slope of this relationship ($M_{pva}$) declined significantly by 13%, thereby indicating reduced PVA at similar LV preload (EDV). Although statistically significant, the associated decrease in the volume intercept, $V_{pva}$, was small and probably of no major physiologic importance. PVA, however, is only an intermediary in the transfer of energy from myocardial O$_2$ consumption (MV O$_2$) to external mechanical work. The efficiency of energy transfer in the final step of myocardial energy transfer from PVA to SW (SW/PVA) fell by 14% (at matched EDV) after the chordae were severed. SW/PVA correlates inversely with the $E_a/E_s$ ratio, and can be approximated by the equation (15): $SW/PVA = 1/([1 + (0.5E_a/E_s)])$. The overall cardiac mechanical efficiency of converting O$_2$ to useful external work (SW/MV O$_2$), however, depends not only on SW/PVA, but also on PVA/MV O$_2$. With a decline in $E_a$ after chordal division, one might anticipate a downward shift of the MV O$_2$–PVA relation to reflect decreased oxygen demand for excitation–contraction coupling (27), thereby increasing PVA/MV O$_2$ efficiency at any PVA while maintaining a relatively constant PVA/excess MV O$_2$ efficiency. This is unlikely, however, since one would not expect chordal division to alter the metabolic state of cardiac myocytes. Moreover, there is clinical evidence that in patients with moderate heart failure mechanical efficiency is reduced, although not to the extent as SW/PVA, presumably due to an increase in the PVA/MV O$_2$ relation (28). In contrast, in patients with severe heart failure who can no longer maintain adequate forward SW, there is a trend towards further reduction in SW/MV O$_2$ due to a substantial fall in the SW/PVA efficiency quotient. In either case, a decline of SW/MV O$_2$ efficiency should accompany chordal division.

Recently, several studies have provided evidence that the left ventricle and the arterial circulation interact in such a manner as to optimize cardiac energetics (28–36), although whether external work (SW) or mechanical efficiency ($SW/MVO_2$) is the critical factor is still a matter of debate. From the standpoint of O$_2$ delivery to the peripheral tissue, optimization of SW would be preferable, whereas optimization of efficiency is more appropriate in terms of cardiac energy expenditure. Based on data obtained from isolated canine heart experiments, Burkhoff and Sagawa (35) developed an analytical model relating LV energetics in terms of ventricular and arterial elastances. Corroborating the work of Sunagawa et al. (31), $SW$ was maximal when $E_a$ equaled $E_s$; however, $SW/MVO_2$ was always maximal when $E_a$ was approximately one-half of $E_s$ at any given $EDV$ (35). In this study, the $E_a/E_s$ ratio was near unity at baseline (0.91 ± 0.53), suggesting that the ventricle and arterial properties were matched such that external stroke work was maximized; however, $E_a/E_s$ was not statistically different from 0.5. Thus, it was also possible that optimal SW/MVO$_2$ might have been the initial operating point during baseline conditions in this experiment. Although $E_a$ fell significantly by 43% after chordal division, $E_a/E_s$ increased due to a proportionally greater (~72%) reduction in $E_s$, indicating that the coupling between pump and load was altered such that neither SW or SW/MVO$_2$ efficiency was optimal. Such is a characteristic of LV pump failure, as demonstrated in dog studies (34) and in patients with failing hearts (28). The effective arterial elastance, as defined by Sunagawa (37), has the dimension of volume elastance (i.e., mmHg/ml). It depends not only on the material properties of the arterial tree, but also on the systemic arterial pressure during ejection. As defined, $E_a$ reflects the ratio of $P_a$ to $SV$ (37). After chordal division, $P_a$ was reduced substantially, as was mean ejection pressure, without any significant change in stroke volume, thereby resulting in a decrease in $E_a$. Although only possibly significant statistically, derived total peripheral resistance fell (as undetectable changes in cardiac output occurred after chordal severing), which was probably chiefly responsible for the decline in $E_a$. Whether this was mediated through direct peripheral beta adrenergic stimulation or via other mechanical reflex responses is not known. It is also possible that the significant, albeit small, decrease in heart rate could have contributed to the fall in $E_a$ since effective arterial elastance is inversely related to the cardiac cycle length (37). This is unlikely, however, given the negligible change (6 min$^{-1}$) in heart rate. The physiologic reason for the decline in $E_a$ is not understood. Since ventriculo–arterial coupling was initially set such that external pressure–volume work was maximal during baseline conditions (i.e., $E_a/E_s \approx 1$), it was possible that in the setting of acute chordal division the lower $E_a$ might reflect some feedback mechanism by which adequate forward cardiac flow output was maintained during moderate heart failure (28). Reduced $E_a$ accompanying LV failure also implies that the ventricle operates more as a pressure pump (small changes in pressure affect a proportionally larger change in stroke volume) since conservation of adequate systemic pressure takes precedence over systemic flow output; this interplay may thereby limit the extent of any potential compensatory decrease in $E_a$. The absolute value of SW/MVO$_2$ efficiency after chordal division, however, might have been compensated for by the observed increase in LV EDV (7), which has a direct, sigmoidal relation to ventricular efficiency (35).
Although the mechanism responsible for valvular–ventricular interaction remains unknown, it should be emphasized that the deterioration of LV systolic mechanics after chordal division, as reflected by declines in these relatively load-independent indices of LV performance, was not due to a change in the intrinsic myocardial contractile state per se, as no myocardial injury occurred. More likely, the decline in LV systolic function resulted from perturbation of the normal three-dimensional geometry of the ventricle. Despite increases in EDD in all three axes after chordal division, the LV chamber became more elliptical at end-diastole. This was due to a relatively greater increment in the LV major axis, as previously reported (38, 39). The small, albeit insignificant, increase in stroke volume after chordal severing, however, was due primarily to augmented fractional dimensional shortening along the two minor LV dimensions. Thus, this increase in chamber eccentricity was possibly present ($P = 0.067$) also at end-ejection. Previously, Rushmer had observed sphericalization of the canine heart in early systole during isovolumic contraction with an abrupt shortening of the major axis and expansion of the minor axes (40). He attributed this phenomenon to the earlier activation and contraction of the papillary muscles, thereby resulting in the descent of the mitral annulus toward the ventricular apex. This initial stretch of the circumferential LV midwall fibers was postulated to enhance LV systolic performance via the Frank-Starling mechanism. We, therefore, hypothesize that the changes in LV geometry associated with disruption of the mitral subvalvular apparatus represents a form of mechanical disadvantage, which leads to the deterioration in LV systolic mechanics and reduced $SW/PA$ efficiency. To our knowledge, this is the first demonstration that a change in LV geometry alone without an alteration in myocardial contractile state can have deleterious effects on LV energetics and ventriculo–arterial coupling.

Several potential limitations of this analysis should be discussed. First, in an attempt to minimize the possibility of overestimating $PA$ (and thereby underestimating $SW/PA$), $V_0$ derived from the quadratic fit of the ESPVR data was used to calculate $PA$. The volume axis intercept originally used by Suga, however, was that obtained directly from measurement of the linear ESPVR (24–26). Since it was conceivable that our conclusions may have been different if PVA had been computed according to Suga’s original definition, LV energetic analysis was repeated using the $V_0$ values derived from the linear ESPVR. Although the absolute values differed, the general results were similar: After the chordae were severed, the slope of the $PA$–EDV relationship declined by 6% ($124±10$ vs. $116±15$ mmHg, $P = 0.024$) with a leftward shift of $V_{pa}$ ($13.7±5.6$ vs. $6.8±6.1$ ml, $P = 0.001$). Similarly, $SW/PA$ at matched LV EDV decreased with chordal division ($0.60±0.14$ vs. $0.38±0.06$, $P = 0.001$). The magnitude of this average decline, however, was substantially greater than that computed from the curvilinear $V_0$ analysis ($35.1±12.1$ for the linear model vs. $13.7±11.4$% for nonlinear, $P < 0.001$). This is consistent with the hypothesis that linearly extrapolated $V_0$ can shift into the negative volume range with chordal severing, thereby artifactualy exaggerating $PA$ and underestimating $SW/PA$. Second, autonomic reflexes might have contributed to the changes observed in this study (despite the theoretical scientific merit of autonomic blockade, we felt this was not realistic in the immediate postcardiopulmonary bypass setting). This was unlikely, however, since both heart rate and total peripheral impedance did not increase despite the deterioration in LV performance after chordal division. This was probably due to the high adrenergic stimulation associated with cardiopulmonary bypass resulting from catecholamine release even though the animals were anesthetized (41). It is possible that the small decrease in heart rate may even reflect some vagal stimulation that could contribute to the deterioration of LV systolic function after chordal division. Again, the physiologic significance of this is unclear given such a negligible change. Third, although we have demonstrated a change in global LV chamber shape, the effect of chordal division on local wall geometry was not assessed. A final potential limitation is the use of a single, steady-state cardiac cycle to estimate $E_a$. Although instantaneous ventriculo–vascular matching can be more accurately determined by computing $E_a$ for each beat during caval occlusion, this measurement may not actually represent actual effective arterial impedance (37). Furthermore, the steady-state ventriculo–arterial interaction is usually considered to have more clinical relevance.

In summary, the current analysis substantiates the validity of the concept of valvular–ventricular interaction. Chordal division resulted in deterioration of LV systolic performance and an increase in the $E_a/E_L$ ratio (because of a proportionally greater decline in $E_L$ compared with $E_a$). This in turn led to a fall in the $SW/PA$ (and, probably, $SW/MVO_2$) efficiency. The integrity of the mitral subvalvular apparatus is, therefore, important not only in terms of global LV pump function, but also in terms of optimization of LV energetics via matching of the left ventricle to the effective arterial load. Although the actual mechanisms responsible for valvular–ventricular interaction have not been elucidated, they may be related to altered three-dimensional LV geometry, and systolic synergy of contraction. These data provide further support to the continuing clinical efforts to preserve as much of the subvalvular structures as possible during mitral valve surgery.

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