Predictors of mitral annulus early diastolic velocity: impact of long-axis function, ventricular filling pattern, and relaxation

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Introduction

Early diastolic velocity (E’) of the mitral annulus used either as an isolated parameter, as the ratio between early (E) velocity of mitral inflow and E’ (E/E’ ratio),1 or in a classification algorithm,2 predicts survival in patients with heart failure (HF) or after myocardial infarction. An accepted explanation of these observations is that, in contrast to the E-wave of mitral inflow, E’ is less sensitive to preload, making it a marker of left ventricular (LV) relaxation.3–6 However, a factor that influences E’ is LV long-axis systolic function, as the length of the downward path of the mitral annulus of the diastole must be identical to its upward path during systole.7 An indirect proof of this proposition is that E’ and peak systolic annular velocity scale symmetrically with the heart size.8

In this retrospective, cross-sectional, observational study we explored the relationship between E’ and LV relaxation, systolic long-axis function and global ventricular filling pattern in three groups of subjects: patients with severe hypertrophic obstructive cardiomyopathy (HOCM), acutely decompressed patients with severe systolic HF, and healthy volunteers. Our hypotheses were that: (1) long-axis systolic function and global LV filling pattern are strongly associated with E’, and (2) that E’ is correlated with LV relaxation parameters even after accounting for systolic long-axis function and global LV filling pattern in each individual group, and in all subjects analysed jointly.

Aims

Although left ventricular (LV) relaxation is well recognized as a predictor of mitral annulus (MA) early diastolic (E’) velocity, its significance relative to other predictors of E’ is less well understood.

Methods and results

We assessed 40 healthy volunteers, 43 patients with acutely decomposed chronic systolic heart failure (HF), and 36 patients with hypertrophic obstructive cardiomyopathy (HOCM) using echocardiography and right or left heart catheterization. Data were obtained at baseline. In addition, in healthy volunteers haemodynamics were varied by graded saline infusion and low body negative pressure, while in HF patients it was varied by vasoactive drug treatment. E- and A-wave velocity (E/A) ratio of the mitral valve inflow, systolic MA velocity integral (s’ integral) and E’ and late velocity (A’) of lateral and septal MA pulsed wave velocities were assessed by echocardiography. Time constant of isovolumic pressure decay (τ₀) was calculated from isovolumic relaxation time/[ln(aortic dicrotic notch pressure) – ln(LV filling pressure)]. In all three groups, s’ integral was the strongest predictor of E’ (partial r = 0.53–0.79; 0.81 for three groups combined), followed by E/A ratio (partial r = 0.10–0.78; 0.26 for all groups combined) and τ₀ (partial r = −0.1 to 0.023; −0.21 for all groups combined).

Conclusion

In healthy adults, patients with systolic HF, or patients with HOCM, E’ is related to LV long-axis function and E/A ratio, a global marker of LV filling. E’ appears less sensitive to LV relaxation.

Keywords

Echocardiography • Relaxation • Tissue Doppler • Heart failure

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Methods

Patients

The study population consisted of 119 subjects stratified into Group 1 (healthy volunteers), Group 2 (systolic HF patients), or Group 3 (HOCM patients), as described below. The rationale of studying these specific groups is that, although the basic relationships between diastolic haemodynamic and echocardiography parameters should be independent of underlying pathology, it appears that the strength of these relationships is modulated by changes in the LV ejection fraction9 and presence of concentric LVH.10

All subjects satisfied the following inclusion criteria: (i) normal sinus rhythm, atrial sensed-ventricular paced rhythm, or atrial and ventricular paced rhythm, (ii) discernible E- and A-waves on the mitral inflow, (iii) no previous mitral valve surgery, or severe mitral regurgitation according to American College of Cardiology/American Heart Association guidelines, (iv) spontaneous ventilation, and (v) no cardiac transplantation.

Group 1 consisted of healthy volunteers who participated in a study that combined invasive and Doppler echocardiography measures to assess the effects of ageing on LV early diastolic function.11 It was composed of 15-young healthy volunteers <50 year (mean age 35 ± 9 years; 12 men) and 25 healthy seniors >65 year (mean age 69 ± 4 year; 13 men). Inclusion criteria and haemodynamic protocol have already been published.11,12 Briefly, all subjects were screened for the presence of arterial hypertension, obstructive coronary artery disease, or structural heart disease, using 24-h blood pressure recordings and baseline and exercise ECGs and echocardiograms. Subjects were excluded if any of the following were present: mean daytime blood pressure >140/90 mmHg, ECG changes suggestive of ischaemic heart disease, left bundle branch block, atrial flutter/fibrillation, atrioventricular block greater than first degree, baseline or exercise-induced wall motion abnormalities, valvular heart disease other than mild valvular insufficiency, right ventricular or LV hypertrophy, untreated thyroid disorders, chronic lung disease, regular cigarette smoking within the previous 10 year, body mass index of ≥30, or use of any cardiovascular medications (e.g., β-blockers, etc.). The Cleveland Clinic Institutional Review Board approved this research project, and informed consent was obtained in all subjects.

Group 2 consisted of 43 consecutive patients >18 year of age with chronic (>6 months) symptomatic systolic HF who underwent Doppler echocardiography and right heart catheterization for optimal evaluation of haemodynamic derangements at the Cleveland Clinic Heart Failure intensive care unit between September 2006 and February 2008. All patients had markedly impaired systolic LV function (defined by the LV ejection fraction 30%) and New York Heart Association class III–IV symptoms. The Cleveland Clinic Institutional Review Board approved this research project, and informed consent was obtained in all subjects.

Finally, Group 3 consisted of 36 consecutive patients, referred for consultation to our institution with suspected obstructive HOCM between October 2006 and April 2008. HOCM was defined by a hypertrophied and non-dilated left ventricle in the absence of another cardiac or systemic disease that could produce a similar magnitude of hypertrophy.13–15 All patients were deemed to have significant dynamic LV outflow tract (LVOT) obstruction that was intractable to maximal medical therapy and were being considered for surgical myectomy vs. alcohol septal ablation. All patients underwent Doppler echocardiography and invasive angiography. The patients are part of a registry that is approved by Cleveland Clinic Institutional Review Board with waiver of individual informed consent.

Sample size was determined by assuming that a sample size of each of the three groups should have a single-sided power of at least 80% to detect a correlation between E′ and relaxation parameter with an absolute r-value of >0.40 or more at the two-tailed alpha level of 0.05. The r-value of 0.40 was selected as a previous study showed an r-value of −0.46 for the correlation between E′ and time constant of the isovolumic pressure decay (τe).6

Invasive and non-invasive data acquisition

Group 1

A 6-Fr balloon-tipped fluid-filled catheter was placed under fluoroscopy through an antecubital vein into the pulmonary artery. Mean pulmonary capillary wedge pressure (PCWP) was determined at end-expiration. Arterial systolic and diastolic blood pressures were measured by cuff sphygmomanometry. After collecting initial haemodynamic and echocardiography data, lower body negative pressure was applied to decrease PCWP as previously reported.13 Briefly, the subject was placed in a Plexiglas box sealed at the level of the iliac crest. The suction was provided with the use of a vacuum pump with a variable autotransformer. Haemodynamic and echocardiography measurements were made after 5 min of LBNP at −15 and −30 mmHg. The negative pressure was then released. After repeated baseline measurements to confirm a return to haemodynamic steady state, cardiac filling was increased by rapid infusion (100 mL/min) of warm (37°C) isotonie saline. Measurements were repeated, after 10 and 20 mL/kg (elderly) or 15 and 30 mL/kg (young) had been infused, to attain similar increases in PCWP in both groups. Complete two-dimensional (2D) and Doppler echocardiography studies were performed, using ATL (Advanced Technology Laboratories; Bothell, Washington) HDI 5000CV. Apical two- and four-chamber views, PW Doppler of inflow and outflow LV tracts, and PW tissue Doppler of lateral and septal side of the mitral annulus were performed at each stage of load protocols.

Group 2

Haemodynamic and echocardiography data were simultaneously collected at baseline (within 12 h of admission) and at follow-up (after 48 h of intensive medical therapy) if the pulmonary artery catheter was still in place. Haemodynamic data, including invasively obtained radial artery blood pressure and PCWP (wedge position was verified by fluoroscopy and phasic changes in pressure waveforms), represent the average of five cycles. Arterial pressures and PCWP were assessed at end expiration. Complete 2D and Doppler echocardiography studies were performed using commercially available equipment. Apical two- and four-chamber views, pulsed-wave PW Doppler interrogation of inflow and outflow LV tracts, and PW tissue Doppler of lateral and septal side of the mitral annulus were performed.

Group 3

Invasive haemodynamics in this group were obtained from a left heart catheterization performed within a mean of 8 ± 9 days of the echocardiography, without change of oral therapy or hospitalization for HF during this period. LV end-diastolic pressures and arterial systolic and diastolic pressures were determined by the catheter placed in the mid-ventricle and ascending aorta, respectively. All patients underwent complete 2D and Doppler echocardiography study that included standard and colour apical two-, three-, four-, and five-chamber views, PW and continuous wave Doppler interrogation of inflow and outflow LV tracts, and PW tissue Doppler of lateral and septal side of the mitral annulus.
In addition, resting LVOT peak velocity was measured by continuous-wave Doppler echocardiography, and resting LVOT pressure gradient was estimated by using simplified Bernoulli equation. Care was taken to avoid contamination of the LVOT waveform by the mitral regurgitation jet. In patients with resting LVOT gradients <30 mmHg, provocative manoeuvres, including Valsalva and amyl nitrate were also used to measure a provokable LVOT gradient. Maximal LVOT gradient was defined as highest recorded gradient (resting or provokable) in a patient. Resting mitral inflow and mitral annulus Doppler data were acquired prior to applying Valsalva manoeuvre and/or amyl nitrate inhalation.

Data analysis
We performed following echocardiography measurements: LV volumes and ejection fraction (Simpson’s biplane rule), LV mass, peak E and late atrial (A) wave velocities of mitral inflow and their ratio, isovolumic relaxation time, duration of E- and A-waves, and E-wave deceleration time. Additionally, we measured s’ wave integral, peak E’ and A’ velocities, and duration of E’ and A’ waves from pulsed wave tissue Doppler tracings of the septal and lateral side of the mitral annulus obtained in the four chamber apical view. For the purpose of analysis, tissue Doppler data obtained from lateral and septal side were averaged.

The time constant of isovolumic pressure decay with a zero asymptote assumption (τ₀) was estimated by the equation:

\[ \tau_0 = \frac{\text{IVRT}}{\ln(AoP_{\text{A0}}) - \ln(\text{filling pressure})} \]

where AoP_{\text{A0}} is aortic dicrotic notch (i.e. end-ejection) pressure and IVRT is isovolumic relaxation time by echocardiography. Aortic dicrotic notch pressure was by the following equation:

\[ AoP_{\text{A0}} = 1.32 \times P_{\text{sys}} + (P_{\text{sys}} - P_{\text{diast}}) \]

where \( P_{\text{sys}} \) is systolic, and \( P_{\text{diast}} \) is diastolic arterial pressure.

Owing to differences in initial protocols, aortic end-ejection pressure and LV filling pressure were calculated slightly differently in three patient groups. In Group 1, arterial pressures were obtained by cuff sphygmomanometer, while LF filling pressures were again represented by PCWP. In Group 2, arterial pressures were obtained from radial arterial pressure tracings, while LF filling pressure was assumed to be represented by PCWP. Finally, in Group 3, systolic and diastolic arterial pressures from aortic blood pressure tracings while LF filling pressure was assumed to be identical to LV end-diastolic pressure.

Statistical methods
Data are shown as mean ± SD unless otherwise noted. If data distribution deviated from normal distribution, appropriate transformations were performed prior to statistical analysis. Comparison between groups was performed by one-way analysis of variance followed by post hoc Tukey’s HSD test, if appropriate.

To assess inter-observer variability of Doppler velocities and time intervals used in the study, two blinded observers measured data of 30 randomly selected patients. To assess intra-observer variability, one observer repeated measurements >1 month after initial measurements. Variability was calculated as the percent variability of the absolute difference between two measurements and expressed as mean ± SD.

To assess if relaxation (i.e. τ₀) affects E’ after controlling for other parameters of LV function, we developed an E’ prediction model based solely on LV systolic long-axis function and its filling pattern. We posit three propositions: (i) systolic ascent of the mitral annulus [i.e. integral of its systolic (s’)] wave is equal to the diastolic descent (i.e. sum of the integrals of the E’ and A’ waves); (ii) the profile of the mitral inflow is similar to the profile of the diastolic mitral annular velocities; and (iii) as E- and A-waves have triangular shape, the mitral inflow profile can be represented by E- and A-wave duration and by the ratio of their velocities (E/A ratio) (see Figure 1). From these propositions, simple algebraic substitutions result in the following equation for the predicted E’ (see Supplementary data online):

\[ \text{Predicted E’} = \frac{E/A}{E/A + A_0/\tau_0} \cdot s’ \]

where E_0 and A_0 are the duration of the E- and A-wave of the mitral inflow, E/A is the ratio of the mitral inflow E- and A-waves, and \( s’ \) is s’ wave integral. Note that E’ predicted in this way is related to LV relaxation only through its effect on E/A ratio. To assess if relaxation has any incremental contribution as an E’ predictor, we then performed a stepwise multiple regression, using predicted E’ and \( \tau_0 \) as predictors of E’.

The null hypothesis was that adding \( \tau_0 \) to predicted E’ would not significantly improve correlation with measured E’.

We also performed stepwise multiple linear regression with E’ as dependent variable, and \( s’ \) integral, E/A ratio and \( \tau_0 \) as predictors. To test if linear regression assumptions were satisfied, we assessed linearity of the plot of observed vs. predicted value (for linearity), the shape of residuals vs. predicted value (for homoscedascity), and tested the normality of residuals distribution. Finally, to assess the impact of multiple measures in individual patients, we applied mixed linear models approach using SPSS 11.0 (IBM Corporation, Somers, NY, USA).

All analyses were performed on whole data sets, as well as on individual subject groups. P-value <0.05 was considered statistically significant.

Results
Table 1 shows baseline clinical and echocardiography variables in our patients. As expected, LV volumes were larger in patients with systolic HF. Again expectedly, LV mass was higher in Groups 2 and 3. In total, the numbers of data-points were as follows: 188 in Group 1, 69 in Group 2, and 36 in Group 3.

There was a moderately strong correlation between mitral inflow E/A ratio and the E’/A’ ratio of the mitral annulus, (r = 0.60, P < 0.0001) as well as between \( s’ \) integral and mitral annulus E’ (r = 0.74, P < 0.0001) (Figure 2A and 2B). The correlation between mitral inflow E/A ratio and the E’/A’ ratio of the mitral annulus was lowest in the SHF group (r = 0.43, P = 0.0002) and highest in the HOCM group (r = 0.83, P < 0.0001). Also, the duration of early diastolic wave of the mitral inflow and of the mitral annulus (E and E’, respectively) strongly correlated (r = 0.89, P < 0.0001), with E-wave having a duration longer by 12 ± 26 ms (P < 0.0001). Similarly, duration of atrial wave of the mitral inflow and of the mitral annulus (A and A’, respectively) correlated (r = 0.75, P < 0.0001), but now with A-wave having a duration slightly shorter by –4 ± 21 ms (P = 0.001). Correlation between mitral annulus E’ and \( \tau_0 \) was significant, but weak (r = –0.19, P = 0.0005; Figure 2C).

There was a strong correlation between the measured and predicted E’ (r = 0.90, P < 0.0001; Figure 3). This correlation was marginally improved by adding \( \tau_0 \) as a predictor (multivariate r = 0.90;
When we repeated this analysis after separating the subjects into groups, \( t_0 \) was a weak independent predictor of \( E' \) in normal subjects (partial \( r^2 = 0.23, P = 0.001 \)) but not in Group 2 (systolic HF patients), or Group 3 (HOCM patients) (Table 2).

Empirical approach to assessment of \( E' \) predictors by multiple linear regression showed that the best overall predictor of \( E' \) was \( s' \) integral (\( P < 0.0001 \)), followed by \( E/A \) ratio (\( P < 0.0001 \)), with \( t_0 \) adding significantly but weakly to the model (partial \( r = -0.21, P = 0.0002 \)). A mixed model that accounted for repeated
measurements of the same subjects confirmed that the strongest $E'$ predictor was $s'$ integral ($P < 0.0001$), followed by $E/A$ ratio ($P < 0.0001$), and $\tau_0$ ($P = 0.0014$). We then repeated this multiple linear regression approach in individual groups of subjects. In contrast to findings described in a previous paragraph, $\tau_0$ showed only a trend towards being a predictor of $E'$ in normal subjects (partial $r = -0.10$, $p = 0.16$). Again, $\tau_0$ was not a predictor of $E'$ in Group 2 (systolic HF patients) or Group 3 (HOCM patients) (Table 3).

**Inter- and intra-observer variability**

The inter-observer variability for mitral valve $E$ and $A$, and mitral annulus $E'$ and $A'$ velocities was $5 \pm 6$, $6 \pm 7$, $6 \pm 8$, and $7 \pm 7\%$, respectively. The inter-observer variability for $s'$ wave integral, mitral $E$ and $A$ duration, and IVRT was $4 \pm 5$, $7 \pm 5$, $8 \pm 6$, and $10 \pm 9\%$, respectively. Intra-observer variabilities were slightly lower (mitral valve $E$ and $A$, and mitral annulus $E'$ and $A'$ velocities were $4 \pm 6$, $6 \pm 6$, $6 \pm 7$, and $7 \pm 7\%$, respectively; $s'$ wave integral, mitral $E$ and $A$ duration, and IVRT was $3 \pm 5$, $6 \pm 5$, $8 \pm 7$, and $9 \pm 9\%$, respectively).

**Uncertainty analysis**

The largest potential source of error in our study is the accuracy of our $\tau_0$ measurements. We have shown that our measurement of $\tau_0$ correlates strongly with $\tau_0$ measured using both Weiss and shifting.

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**Figure 2** (A) Correlation between the ratio of the peak early and late diastolic mitral inflow velocities ($E/A$ ratio) and the ratio of the peak early and late diastolic velocities of the mitral annulus ($E'/A'$ ratio); (B) correlation between peak early diastolic velocity of the mitral annulus ($E'$) and the integral of the systolic velocity of the mitral annulus ($s'$ integral); (C) correlation between peak early diastolic velocity of the mitral annulus ($E'$) and the time constant of the isovolumic pressure decay ($\tau_0$). HOCM, hypertrophic obstructive cardiomyopathy; SHF, systolic heart failure; Elderly, elderly healthy volunteers; Young, young healthy volunteers.

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asymptote method. However, to further estimate the maximal potential correlation between \( \tau_0 \) and \( E' \) after adjusting for \( s' \) and \( E' \) ratio, we first subtracted the variance attributable to \( E' \) measurement from the residual sum of squares of multiple linear regression of \( E' \) predictors. Assuming that \( \tau_0 \) explained the entire residual sum of squares after this subtraction, its partial correlation coefficient was only \(-0.47\), a value still not high enough for \( E' \) to be a clinically useful surrogate for \( \tau_0 \).

**Discussion**

In this paper, we analyse the way LV long-axis function, filling pattern, and relaxation relate to \( E' \), a marker of diastolic function. Partial correlations we obtained by multiple regression indicate that the dominant factors associated with \( E' \), in three groups tested either separately or jointly, were the LV long-axis function followed by LV filling pattern, with LV relaxation having a smaller impact. These findings are further strengthened by showing that adding LV relaxation to \( E' \) calculated by a mathematical equation that incorporates LV long-axis function and filling pattern only minimally improves accuracy of \( E' \) prediction.

\[ y = 0.9x + 1.2 \]
\[ r = 0.90 \]
\[ p < 0.001 \]
\[ n = 293 \]

![Figure 3](image-url) Correlation between measured peak early diastolic velocity of the mitral annulus (\( E' \)) and \( E' \) predicted from the systolic displacement of the mitral annulus and the profile of the mitral inflow (predicted \( E' \)).

**Table 2** Estimating early diastolic velocity of the mitral annulus (\( E' \)) from predicted \( E' \) and time constant of the isovolumic pressure decay (\( \tau_0 \)) by multiple linear regression

<table>
<thead>
<tr>
<th></th>
<th>( r, ) full model (95% CI)</th>
<th>Predicted ( E' ) partial ( r ) (95% CI)</th>
<th>( P )-value</th>
<th>( \tau_0 ) Partial ( r ) (95% CI)</th>
<th>( P )-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All subjects</td>
<td>0.90 (0.88–0.92)</td>
<td>0.90 (0.88–0.92)</td>
<td>&lt;0.0001</td>
<td>-0.13 (-0.24 to -0.02)</td>
<td>0.03</td>
</tr>
<tr>
<td>HOCM</td>
<td>0.83 (0.69–0.91)</td>
<td>0.83 (0.69–0.91)</td>
<td>&lt;0.0001</td>
<td>0.01 (-0.32 to 0.34)</td>
<td>0.99</td>
</tr>
<tr>
<td>SHF</td>
<td>0.54 (0.35–0.69)</td>
<td>0.54 (0.35–0.69)</td>
<td>&lt;0.0001</td>
<td>-0.064 (-0.30 to 0.18)</td>
<td>0.6</td>
</tr>
<tr>
<td>Healthy volunteers</td>
<td>0.86 (0.82–0.89)</td>
<td>0.86 (0.82–0.89)</td>
<td>&lt;0.0001</td>
<td>-0.23 (-0.36 to -0.09)</td>
<td>0.001</td>
</tr>
</tbody>
</table>

CI, confidence interval; HOCM, hypertrophic cardiomyopathy; SHF, systolic heart failure.

**\( E' \) as a predictor of relaxation**

Several studies assessed \( E' \) as an estimate of invasively quantified LV relaxation. Nagueh et al. studied 10 dogs that were subjected to various haemodynamic perturbations to show a strong relationship between \( E' \) and \( \tau \) with an \( r \) of \(-0.83\). In a similar study, Firstenberg et al. showed that, while \( E' \) and \( \tau \) correlated with an \( r \) of \(-0.70\), \( E' \) and E-wave velocities changed by a similar percentage in the setting of an isolated change in preload, indicating that \( E' \) is preload dependent. In contrast, clinical studies that assessed the same issue came up with lower correlations. Ommen et al. showed in a study of 100 subjects a correlation of \( r \) of \(-0.46\). Similarly, in study of 12 control patients and 43 subjects with HF and normal ejection fraction, Kasner et al. obtained an \( r \) of \(-0.33\), while Mizuno et al. in a study of 117 patients showed a correlation of \(-0.40\) (using a standard zero-asymptote method for \( \tau \) measurement). Our study in 119 subjects shows a correlation between \( E' \) and \( \tau \) that, when compared with these studies, was even lower, although with confidence intervals approaching the values reported before (\( r = -0.19 \), 95% CI: \(-0.01 \) to \(-0.36\)). The reason for lower correlation values obtained in this study may be multifactorial. While previously validated, our method of calculating \( \tau \) should be viewed as an estimation. Additionally, some studies suggest that relaxation more strongly predicts \( E' \) in the setting of decreased systolic function, and as most of our subject had normal ejection fractions, our findings may be biased. However, in our study \( \tau_0 \) was not a strong predictor of relaxation even in patients with systolic HF.

**Relationship between left ventricular long-axis systolic displacement and early diastolic myocardial velocity**

Conservation of mass necessitates that systolic and diastolic displacements are identical, which makes systolic and diastolic velocities interdependent. Indeed, Mogelvang et al. showed a correlation between \( s' \) and \( E' \) velocity of \( r = 0.52 \) even after adjustment for age, sex, body mass index, and heart rate. Furthermore, \( s' \) and \( E' \) velocity decrease in parallel in the systolic and diastolic HF.23 We extend these findings by showing that \( E' \) velocity strongly depends on the \( s' \) integral, a measure of systolic displacement. We furthermore show that our predicted \( E' \), which is very much determined by systolic displacement, efficiently tracks true \( E' \) values.
Left ventricular filling pattern and the local profiles of myocardial diastolic velocities

LV filling pattern in sinus rhythm has two components: E-wave of early, and A-wave of late filling. The determinants of LV filling are well-known, with the interaction of relaxation and ventricular stiffness determining early, and atrial contractility and load determining late filling. LV filling pattern, by default, is the product of local ventricular behaviour. Given that myocardial structure throughout ventricle carries some amount of homogeneity one can surmise that this homogeneity of structure translates to some homogeneity of function (i.e. local relaxation and stiffness parameters), resulting in regional myocardial velocities having at least some between-region similarity. Indeed, we do show that E/A ratios of mitral inflow and long-axis velocities, while not identical, are similar.

Clinical implications

The echocardiographic detection of LV diastolic dysfunction has been made more difficult by the confusing terminology, including the question of what exactly constitutes diastolic dysfunction. Elements of diastolic function, such as active relaxation and chamber compliance that are traditionally measured using invasive pressure and volume data, do not have direct echocardiographic counterparts. Rather, echocardiographic indices reflect combined influences of relaxation, compliance, and interventricular dependence. As a result, diagnosis of elevated filling pressures by echocardiography is not straightforward. Our findings reinforce the idea that, instead of relying on a single parameter for the assessment of LV diastolic function (or filling pressures), several parameters should be combined.\(^7\) Of note, while decision-making algorithms were proposed by leading international echocardiography associations,\(^8\) there is still a paucity of data how well they perform in the clinical setting, especially when discordant individual parameters occur. Additionally, E’ measurement may have prognostic implications even beyond diastolic function assessment.\(^{1,26}\) These issues have to addressed by future studies.

Limitations

This was a retrospective study with slightly different protocols in each of the studied groups. However, sample size of each group was large enough to adequately detect a relevant correlation between E’ and relaxation. Inclusion of patients with mitral regurgitation, paced ventricular rhythms, and severe LV hypertrophy may reduce the correlation between E’ and LV relaxation. Also, while it appears that E’ is less sensitive to LV relaxation than to systolic long-axis function and E/A ratio in patients we studied, it is uncertain if that can be generalized to all cardiovascular states, such as patients with mild to moderate systolic dysfunction, hypertensive heart disease, restrictive cardiomyopathies, or diabetic heart disease.

E’-\(\tau_0\) correlation performed after controlling for E/A values may underestimate intrinsic relationship between E’ and \(\tau_0\). Also, our method of \(\tau_0\) estimation is less sophisticated and may be more prone to errors than non-linear regression methods that are traditionally used in rigorous haemodynamic assessment and was measured slightly differently in three patient groups. Our method is further compromised by lack of simultaneity in echo and invasive measurements in HOCM patients, which may introduce additional measurement error. Because of this, our ability to detect weak E’ correlations with tau after accounting for systolic long-axis function and global LV filling pattern (hypothesis 2). However, weak correlations are less relevant, it has been shown that our \(\tau_0\) estimation method correlates satisfactorily with a standard zero—and shifting—asymptote methods,\(^{11,18}\) and has been previously used in clinical research.\(^{11,27}\)

Another technical limitation is a known presence of measurement error that is inherent in measurement of echocardiography variables, especially in the setting of decreased mitral annulus velocities.

Conclusion

In this paper, we show that E’ is related LV long-axis displacement during systole and that the diastolic pattern of long-axis motion largely follows the mitral inflow pattern. E’ is comparatively less sensitive to changes in LV relaxation in healthy adults, patients with systolic HF, or patients with HOCM.

Supplementary data

Supplementary data are available at European Journal of Echocardiography online.
Conflict of interest: none declared.

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